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(54) Title: METHODS OF DIAGNOSIS OF ANGIOGENESIS, COMPOSITIONS AND METHODS OF SCREENING FOR ANGIOGENESIS MODULATORS

(57) Abstract: Described herein are methods and compositions that can be used for diagnosis and treatment of angiogenic phenotypes and angiogenesis-associated diseases. Also described herein are methods that can be used to identify modulators of angiogenesis.

METHODS OF DIAGNOSIS OF ANGIOGENESIS, COMPOSITIONS AND METHODS OF SCREENING FOR ANGIOGENESIS MODULATORS

CROSS-REFERENCES TO RELATED APPLICATIONS

This application claims priority to USSN 09/784,356, filed February 14 2001; USSN 09/791,390, filed February 22, 2001; USSN 60/285,475, filed April 19, 2001, USSN 60/310,025, filed August 3, 2001, and USSN 60/334,244, filed November 29, 2001, each of which is herein incorporated by reference in its entirety.

FIELD OF THE INVENTION

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The invention relates to the identification of nucleic acid and protein expression profiles and nucleic acids, products, and antibodies thereto that are involved in angiogenesis; and to the use of such expression profiles and compositions in diagnosis and therapy of angiogenesis. The invention further relates to methods for identifying and using agents and/or targets that modulate angiogenesis.

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BACKGROUND OF THE INVENTION

Both vasculogenesis, the development of an interactive vascular system comprising arteries and veins, and angiogenesis, the generation of new blood vessels, play a role in embryonic development. In contrast, angiogenesis is limited in a normal adult to the placenta, ovary, endometrium and sites of wound healing. However, angiogenesis, or its absence, plays an important role in the maintenance of a variety of pathological states. Some of these states are characterized by neovascularization, e.g., cancer, diabetic retinopathy, glaucoma, and age related macular degeneration. Others, e.g., stroke, infertility, heart disease, ulcers, and scleroderma, are diseases of angiogenic insufficiency.

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Angiogenesis has a number of stages (see, e.g., Folkman, J.Natl Cancer Inst. 82:4-6, 1990; Firestein, J Clin Invest. 103:3-4, 1999; Koch, Arthritis Rheum. 41:951-62, 1998; Carter, Oncologist 5(Suppl 1):51-4, 2000; Browder et al., Cancer Res. 60:1878-86, 2000; and Zhu and Witte, Invest New Drugs 17:195-212, 1999). The early stages of angiogenesis

include endothelial cell protease production, migration of cells, and proliferation. The early stages also appear to require some growth factors, with VEGF, TGF-α, angiostatin, and selected chemokines all putatively playing a role. Later stages of angiogenesis include population of the vessels with mural cells (pericytes or smooth muscle cells), basement membrane production, and the induction of vessel bed specializations. The final stages of vessel formation include what is known as "remodeling", wherein a forming vasculature becomes a stable, mature vessel bed. Thus, the process is highly dynamic, often requiring coordinated spatial and temporal waves of gene expression.

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Conversely, the complex process may be subject to disruption by interfering with one or more critical steps. Thus, the lack of understanding of the dynamics of angiogenesis prevents therapeutic intervention in serious diseases such as those indicated. It is an object of the invention to provide methods that can be used to screen compounds for the ability to modulate angiogenesis. Additionally, it is an object to provide molecular targets for therapeutic intervention in disease states which either have an undesirable excess or a deficit in angiogenesis. The present invention provides solutions to both.

SUMMARY OF THE INVENTION

The present invention provides compositions and methods for detecting or modulating angiogenesis associated sequences.

In one aspect, the invention provides a method of detecting an angiogenesis-associated transcript in a cell in a patient, the method comprising contacting a biological sample from the patient with a polynucleotide that selectively hybridized to a sequence at least 80% identical to a sequence as shown in Tables 1-8. In one embodiment, the biological sample is a tissue sample. In another embodiment, the biological sample comprises isolated nucleic acids, which are often mRNA.

In another embodiment, the method further comprises the step of amplifying nucleic acids before the step of contacting the biological sample with the polynucleotide. Often, the polynucleotide comprises a sequence as shown in Tables 1-8. The polynucleotide can be labeled, for example, with a fluorescent label and can be immobilized on a solid surface.

In other embodiments the patient is undergoing a therapeutic regimen to treat a disease associated with angiogenesis or the patient is suspected of having an angiogenesis-associated disorder.

In another aspect, the invention comprises an isolated nucleic acid molecule consisting of a polynucleotide sequence as shown in Tables 1-8. The nucleic acid molecule can be labeled, for example, with a fluorescent label,

In other aspects, the invention provides an expression vector comprising an isolated nucleic acid molecule consisting of a polynucleotide sequence as shown in Tables 1-8 or a host cell comprising the expression vector.

In another embodiment, the isolated nucleic acid molecule encodes a polypeptide having an amino acid sequence as shown in Table 8.

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In another aspect, the invention provides an isolated polypeptide which is encoded by a nucleic acid molecule having polynucleotide sequence as shown in Tables 1-8. In one embodiment, the isolated polypeptide has an amino acid sequence as shown in Table 8.

In another embodiment, the invention provides an antibody that specifically binds a polypeptide that has an amino acid sequence as shown in Table 8 or which is encoded by a nucleotide sequence of Tables 1-8. The antibody can be conjugated or fused to an effector component such as a fluorescent label, a toxin, or a radioisotope. In some embodiments, the antibody is an antibody fragment or a humanized antibody.

In another aspect, the invention provides a method of detecting a cell undergoing angiogenesis in a biological sample from a patient, the method comprising contacting the biological sample with an antibody that specifically binds to a polypeptide that has an amino acid sequence as shown in Table 8 or which is encoded by a nucleotide sequence of Tables 1-8. In some embodiments, the antibody is further conjugated or fused to an effector component, for example, a fluorescent label.

In another embodiment, the invention provides a method of detecting antibodies specific to angiogenesis in a patient, the method comprising contacting a biological sample from the patient with a polypeptide which is encoded by a nucleotide sequence of Tables 1-8.

The invention also provides a method of identifying a compound that modulates the activity of an angiogenesis-associated polypeptide, the method comprising the steps of: (i) contacting the compound with a polypeptide that comprises at least 80% identity to an amino acid sequence as shown in Table 8 or which is encoded by a nucleotide sequence of Tables 1-8; and (ii) detecting an increase or a decrease in the activity of the polypeptide. In one embodiment, the polypeptide has an amino acid sequence as shown in Table 8 or is a

polypeptide encoded by a nucleotide sequence of Tables 1-8. In another embodiment, the polypeptide is expressed in a cell.

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The invention also provides a method of identifying a compound that modulates angiogenesis, the method comprising steps of: (i) contacting the compound with a cell undergoing angiogenesis; and (ii) detecting an increase or a decrease in the expression of a polypeptide sequence as shown in Table 8 or a polypeptide which is encoded by a nucleotide sequence of Tables 1-8. In one embodiment, the detecting step comprises hybridizing a nucleic acid sample from the cell with a polynucleotide that selectively hybridizes to a sequence at least 80% identical to a sequence as shown in Tables 1-8. In another embodiment, the method further comprises detecting an increase or decrease in the expression of a second sequence as shown in Table 8 or a polypeptide which is encoded by a nucleotide sequence of Tables 1-8.

In another embodiment, the invention provides a method of inhibiting angiogenesis in a cell that expresses a polypeptide at least 80% identical to a sequence as shown in Table 8 or which is 80% identical to a polypeptide encoded by a nucleotide sequence of Tables 1-8, the method comprising the step of contacting the cell with a therapeutically effective amount of an inhibitor of the polypeptide. In one embodiment, the polypeptide has an amino acid sequence shown in Table 8 or is a polypeptide which is encoded by a nucleotide sequence of Tables 1-8. In another embodiment, the inhibitor is an antibody.

In other embodiments, the invention provides a method of activating angiogenesis in a cell that expresses a polypeptide at least 80% identical to a sequence as shown in Table 8 or at least 80% identical to a polypeptide which is encoded by a nucleotide sequence of Tables 1-8, the method comprising the step of contacting the cell with a therapeutically effective amount of an activator of the polypeptide. In one embodiment, the polypeptide has an amino acid sequence shown in Table 8 or is a polypeptide which is encoded by a nucleotide sequence of Tables 1-8.

Other aspects of the invention will become apparent to the skilled artisan by the following description of the invention.

Tables 1-8 provide nucleotide sequence of genes that exhibit changes in expression levels as a function of time in tissue undergoing angiogenesis compared to tissue that is not.

DESCRIPTION OF THE SPECIFIC EMBODIMENTS

In accordance with the objects outlined above, the present invention provides novel methods for diagnosis and treatment of disorders associated with angiogenesis (sometimes referred to herein as angiogenesis disorders or AD), as well as methods for screening for compositions which modulate angiogenesis. By "disorder associated with angiogenesis" or "disease associated with angiogenesis" herein is meant a disease state which is marked by either an excess or a deficit of blood vessel development. Angiogenesis disorders associated with increased angiogenesis include, but are not limited to, cancer and proliferative diabetic retinopathy. Pathological states for which it may be desirable to increase angiogenesis include stroke, heart disease, infertility, ulcers, wound healing, ischemia, and scleradoma. Solid tumors typically require angiogenesis to support or sustain growth, e.g., breast, colon, lung, brain, bladder, and prostate tumors. Other AD include, e.g., arthritis, inflammatory bowel disease, diabetis retinopathy, macular degeneration, atherosclerosis, and psoriasis. Also provided are methods for treating AD.

15 Definitions

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The term "angiogenesis protein" or "angiogenesis polynucleotide" refers to nucleic acid and polypeptide polymorphic variants, alleles, mutants, and interspecies homologs that: (1) have an amino acid sequence that has greater than about 60% amino acid sequence identity, 65%, 70%, 75%, 80%, 85%, 90%, preferably 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98% or 99% or greater amino acid sequence identity, preferably over a region of over a region of at least about 25, 50, 100, 200, 500, 1000, or more amino acids, to an angiogenesis protein sequence of Table 8; (2) bind to antibodies, e.g., polyclonal antibodies, raised against an immunogen comprising an amino acid sequence of Table 8, and conservatively modified variants thereof; (3) specifically hybridize under stringent hybridization conditions to an anti-sense strand corresponding to a nucleic acid sequence of Tables 1-8 and conservatively modified variants thereof; (4) have a nucleic acid sequence that has greater than about 95%, preferably greater than about 96%, 97%, 98%, 99%, or higher nucleotide sequence identity, preferably over a region of at least about 25, 50, 100, 200, 500, 1000, or more nucleotides, to a sense sequence corresponding to one set out in Tables 1-8. A polynucleotide or polypeptide sequence is typically from a mammal including, but not limited to, primate, e.g., human; rodent, e.g., rat, mouse, hamster; cow, pig, horse, sheep, or any mammal. An "angiogenesis polypeptide" and an "angiogenesis polynucleotide," include both naturally occurring or recombinant.

A "full length" angiogenesis protein or nucleic acid refers to an agiogenesis polypeptide or polynucleotide sequence, or a variant thereof, that contains all of the elements normally contained in one or more naturally occurring, wild type angiogenesis polynucleotide or polypeptide sequences. The "full length" may be prior to, or after, various stages of post-translation processing.

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"Biological sample" as used herein is a sample of biological tissue or fluid that contains nucleic acids or polypeptides, e.g., of an angiogenic protein. Such samples include, but are not limited to, tissue isolated from primates, e.g., humans, or rodents, e.g., mice, and rats. Biological samples may also include sections of tissues such as biopsy and autopsy samples, and frozen sections taken for histologic purposes. A biological sample is typically obtained from a eukaryotic organism, most preferably a mammal such as a primate e.g., chimpanzee or human; cow; dog; cat; a rodent, e.g., guinea pig, rat, mouse; rabbit; or a bird; reptile; or fish.

"Providing a biological sample" means to obtain a biological sample for use in methods described in this invention. Most often, this will be done by removing a sample of cells from an animal, but can also be accomplished by using previously isolated cells (e.g., isolated by another person, at another time, and/or for another purpose), or by performing the methods of the invention in vivo. Archival tissues, having treatment or outcome histroy, will be particularly useful.

The terms "identical" or percent "identity," in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same (i.e., about 70% identity, preferably 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99%, or higher identity over a specified region (e.g., SEQ ID NOS:1-229), when compared and aligned for maximum correspondence over a comparison window or designated region) as measured using a BLAST or BLAST 2.0 sequence comparison algorithms with default parameters described below, or by manual alignment and visual inspection (see, e.g., NCBI web site http://www.ncbi.nlm.nih.gov/BLAST/ or the like). Such sequences are then said to be "substantially identical." This definition also refers to, or may be applied to, the compliment of a test sequence. The definition also includes sequences that have deletions and/or additions, as well as those that have substitutions. As described below, the preferred algorithms can account for gaps and the like. Preferably, identity exists over a region that is at least about 25 amino acids or nucleotides in length, or more preferably over a region that is 50-100 amino acids or nucleotides in length.

For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are entered into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. Preferably, default program parameters can be used, or alternative parameters can be designated. The sequence comparison algorithm then calculates the percent sequence identities for the test sequences relative to the reference sequence, based on the program parameters.

A "comparison window", as used herein, includes reference to a segment of any one of the number of contiguous positions selected from the group consisting of from 20 to 600, usually about 50 to about 200, more usually about 100 to about 150 in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. Methods of alignment of sequences for comparison are well-known in the art. Optimal alignment of sequences for comparison can be conducted, e.g., by the local homology algorithm of Smith & Waterman, Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman & Wunsch, J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson & Lipman, Proc. Nat'l. Acad. Sci. USA 85:2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by manual alignment and visual inspection (see, e.g., Current Protocols in Molecular Biology (Ausubel et al., eds. 1995 supplement)).

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A preferred example of algorithm that is suitable for determining percent sequence identity and sequence similarity are the BLAST and BLAST 2.0 algorithms, which are described in Altschul et al., Nuc. Acids Res. 25:3389-3402 (1977) and Altschul et al., J. Mol. Biol. 215:403-410 (1990), respectively. BLAST and BLAST 2.0 are used, with the parameters described herein, to determine percent sequence identity for the nucleic acids and proteins of the invention. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul et al., supra). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are extended in both directions along each sequence for as

far as the cumulative alignment score can be increased. Cumulative scores are calculated using, for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, M=5, N=4 and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength of 3, and expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff, Proc. Natl. Acad. Sci. USA 89:10915 (1989)) alignments (B) of 50, expectation (E) of 10, M=5, N=4, and a comparison of both strands.

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The BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin & Altschul, Proc. Nat'l. Acad. Sci. USA 90:5873-5787 (1993)). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.2, more preferably less than about 0.01, and most preferably less than about 0.001.

An indication that two nucleic acid sequences or polypeptides are substantially identical is that the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the antibodies raised against the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second polypeptide, for example, where the two peptides differ only by conservative substitutions. Another indication that two nucleic acid sequences are substantially identical is that the two molecules or their complements hybridize to each other under stringent conditions, as described below. Yet another indication that two nucleic acid sequences are substantially identical is that the same primers can be used to amplify the sequences.

A "host cell" is a naturally occurring cell or a transformed cell that contains an expression vector and supports the replication or expression of the expression vector. Host cells may be cultured cells, explants, cells *in vivo*, and the like. Host cells may be

prokaryotic cells such as *E. coli*, or eukaryotic cells such as yeast, insect, amphibian, or mammalian cells such as CHO, HeLa, and the like (see, *e.g.*, the American Type Culture Collection catalog or web site, www.atcc.org).

The terms "polypeptide," "peptide" and "protein" are used interchangeably herein to refer to a polymer of amino acid residues. The terms apply to amino acid polymers in which one or more amino acid residue is an artificial chemical mimetic of a corresponding naturally occurring amino acid, as well as to naturally occurring amino acid polymers and non-naturally occurring amino acid polymer.

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The term "amino acid" refers to naturally occurring and synthetic amino acids, as well as amino acid analogs and amino acid mimetics that function in a manner similar to the naturally occurring amino acids. Naturally occurring amino acids are those encoded by the genetic code, as well as those amino acids that are later modified, e.g., hydroxyproline, γ-carboxyglutamate, and O-phosphoserine. Amino acid analogs refers to compounds that have the same basic chemical structure as a naturally occurring amino acid, i.e., an α carbon that is bound to a hydrogen, a carboxyl group, an amino group, and an R group, e.g., homoserine, norleucine, methionine sulfoxide, methionine methyl sulfonium. Such analogs have modified R groups (e.g., norleucine) or modified peptide backbones, but retain the same basic chemical structure as a naturally occurring amino acid. Amino acid mimetics refers to chemical compounds that have a structure that is different from the general chemical structure of an amino acid, but that functions in a manner similar to a naturally occurring amino acid.

Amino acids may be referred to herein by either their commonly known three letter symbols or by the one-letter symbols recommended by the IUPAC-IUB Biochemical Nomenclature Commission. Nucleotides, likewise, may be referred to by their commonly accepted single-letter codes.

"Conservatively modified variants" applies to both amino acid and nucleic acid sequences. With respect to particular nucleic acid sequences, conservatively modified variants refers to those nucleic acids which encode identical or essentially identical amino acid sequences, or where the nucleic acid does not encode an amino acid sequence, to essentially identical sequences. Because of the degeneracy of the genetic code, a large number of functionally identical nucleic acids encode any given protein. For instance, the codons GCA, GCC, GCG and GCU all encode the amino acid alanine. Thus, at every position where an alanine is specified by a codon, the codon can be altered to any of the corresponding codons described without altering the encoded polypeptide. Such nucleic acid variations are "silent variations," which are one species of conservatively modified

variations. Every nucleic acid sequence herein which encodes a polypeptide also describes every possible silent variation of the nucleic acid. One of skill will recognize that each codon in a nucleic acid (except AUG, which is ordinarily the only codon for methionine, and TGG, which is ordinarily the only codon for tryptophan) can be modified to yield a functionally identical molecule. Accordingly, each silent variation of a nucleic acid which encodes a polypeptide is implicit in each described sequence with respect to the expression product, but not with respect to actual probe sequences.

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As to amino acid sequences, one of skill will recognize that individual substitutions, deletions or additions to a nucleic acid, peptide, polypeptide, or protein sequence which alters, adds or deletes a single amino acid or a small percentage of amino acids in the encoded sequence is a "conservatively modified variant" where the alteration results in the substitution of an amino acid with a chemically similar amino acid.

Conservative substitution tables providing functionally similar amino acids are well known in the art. Such conservatively modified variants are in addition to and do not exclude polymorphic variants, interspecies homologs, and alleles of the invention.

The following eight groups each contain amino acids that are conservative substitutions for one another:1) Alanine (A), Glycine (G); 2) Aspartic acid (D), Glutamic acid (E); 3) Asparagine (N), Glutamine (Q); 4) Arginine (R), Lysine (K); 5) Isoleucine (I), Leucine (L), Methionine (M), Valine (V); 6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W); 7) Serine (S), Threonine (T); and 8) Cysteine (C), Methionine (M) (see, e.g., Creighton, Proteins (1984)).

Macromolecular structures such as polypeptide structures can be described in terms of various levels of organization. For a general discussion of this organization, see, e.g., Alberts et al., Molecular Biology of the Cell (3rd ed., 1994) and Cantor and Schimmel, Biophysical Chemistry Part I: The Conformation of Biological Macromolecules (1980). "Primary structure" refers to the amino acid sequence of a particular peptide. "Secondary structure" refers to locally ordered, three dimensional structures within a polypeptide. These structures are commonly known as domains. Domains are portions of a polypeptide that form a compact unit of the polypeptide and are typically 25 to approximately 500 amino acids long. Typical domains are made up of sections of lesser organization such as stretches of β -sheet and α -helices. "Tertiary structure" refers to the complete three dimensional structure of a polypeptide monomer. "Quaternary structure" refers to the three dimensional

structure formed, usually by the noncovalent association of independent tertiary units. Anisotropic terms are also known as energy terms.

A "label" or a "detectable moiety" is a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, chemical, or other physical means. For example, useful labels include ³²P, fluorescent dyes, electron-dense reagents, enzymes (e.g., as commonly used in an ELISA), biotin, digoxigenin, or haptens and proteins which can be made detectable, e.g., by incorporating a radiolabel into the peptide or used to detect antibodies specifically reactive with the peptide.

An "effector" or "effector moiety" or "effector component" is a molecule that is bound (or linked, or conjugated), either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds, to an antibody. The "effector" can be a variety of molecules including, for example, detection moieties including radioactive compounds, fluroescent compounds, an enzyme or substrate, tags such as epitope tags, a toxin; a chemotherapeutic agent; a lipase; an antibiotic; or a radioisotope emitting "hard" e.g., beta radiation.

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A "labeled nucleic acid probe or oligonucleotide" is one that is bound, either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds to a label such that the presence of the probe may be detected by detecting the presence of the label bound to the probe. Alternatively, method using high affinity interactions may achieve the same results where one of a pair of binding partners binds to the other, e.g., biotin, streptavidin.

As used herein a "nucleic acid probe or oligonucleotide" is defined as a nucleic acid capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation. As used herein, a probe may include natural (i.e., A, G, C, or T) or modified bases (7-deazaguanosine, inosine, etc.). In addition, the bases in a probe may be joined by a linkage other than a phosphodiester bond, so long as it does not interfere with hybridization. Thus, for example, probes may be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages. It will be understood by one of skill in the art that probes may bind target sequences lacking complete complementarity with the probe sequence depending upon the stringency of the hybridization conditions. The probes are preferably directly labeled as with isotopes, chromophores, lumiphores, chromogens, or indirectly labeled such as with biotin to which a streptavidin

complex may later bind. By assaying for the presence or absence of the probe, one can detect the presence or absence of the select sequence or subsequence.

The term "recombinant" when used with reference, e.g., to a cell, or nucleic acid, protein, or vector, indicates that the cell, nucleic acid, protein or vector, has been modified by the introduction of a heterologous nucleic acid or protein or the alteration of a native nucleic acid or protein, or that the cell is derived from a cell so modified. Thus, for example, recombinant cells express genes that are not found within the native (non-recombinant) form of the cell or express native genes that are otherwise abnormally expressed, under expressed or not expressed at all.

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The term "heterologous" when used with reference to portions of a nucleic acid indicates that the nucleic acid comprises two or more subsequences that are not found in the same relationship to each other in nature. For instance, the nucleic acid is typically recombinantly produced, having two or more sequences from unrelated genes arranged to make a new functional nucleic acid, e.g., a promoter from one source and a coding region from another source. Similarly, a heterologous protein indicates that the protein comprises two or more subsequences that are not found in the same relationship to each other in nature (e.g., a fusion protein).

A "promoter" is defined as an array of nucleic acid control sequences that direct transcription of a nucleic acid. As used herein, a promoter includes necessary nucleic acid sequences near the start site of transcription, such as, in the case of a polymerase II type promoter, a TATA element. A promoter also optionally includes distal enhancer or repressor elements, which can be located as much as several thousand base pairs from the start site of transcription. A "constitutive" promoter is a promoter that is active under most environmental and developmental conditions. An "inducible" promoter is a promoter that is active under environmental or developmental regulation. The term "operably linked" refers to a functional linkage between a nucleic acid expression control sequence (such as a promoter, or array of transcription factor binding sites) and a second nucleic acid sequence, wherein the expression control sequence directs transcription of the nucleic acid corresponding to the second sequence.

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An "expression vector" is a nucleic acid construct, generated recombinantly or synthetically, with a series of specified nucleic acid elements that permit transcription of a particular nucleic acid in a host cell. The expression vector can be part of a plasmid, virus, or nucleic acid fragment. Typically, the expression vector includes a nucleic acid to be transcribed operably linked to a promoter.

The phrase "selectively (or specifically) hybridizes to" refers to the binding, duplexing, or hybridizing of a molecule only to a particular nucleotide sequence under stringent hybridization conditions when that sequence is present in a complex mixture (e.g., total cellular or library DNA or RNA).

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The phrase "stringent hybridization conditions" refers to conditions under which a probe will hybridize to its target subsequence, typically in a complex mixture of nucleic acids, but to no other sequences. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in Tijssen, Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Probes, "Overview of principles of hybridization and the strategy of nucleic acid assays" (1993). Generally, stringent conditions are selected to be about 5-10°C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength pH. The T_m is the temperature (under defined ionic strength, pH, and nucleic concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at T_m, 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g., 10 to 50 nucleotides) and at least about 60°C for long probes (e.g., greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. For selective or specific hybridization, a positive signal is at least two times background, preferably 10 times background hybridization. Exemplary stringent hybridization conditions can be as following: 50% formamide, 5x SSC, and 1% SDS, incubating at 42°C, or, 5x SSC, 1% SDS, incubating at 65°C, with wash in 0.2x SSC, and 0.1% SDS at 65°C. For PCR, a temperature of about 36°C is typical for low stringency amplification, although annealing temperatures may vary between about 32°C and 48°C depending on primer length. For high stringency PCR amplification, a temperature of about 62°C is typical, although high stringency annealing temperatures can range from about 50°C to about 65°C, depending on the primer length and specificity. Typical cycle conditions for both high and low stringency amplifications include a denaturation phase of 90°C - 95°C for 30 sec - 2 min., an annealing phase lasting 30 sec. - 2 min., and an extension phase of about 72°C for 1 - 2 min. Protocols and guidelines for low and high stringency amplification

reactions are provided, e.g., in Innis et al. (1990) PCR Protocols, A Guide to Methods and Applications, Academic Press, Inc. N.Y.).

Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides which they encode are substantially identical. This occurs, for example, when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code. In such cases, the nucleic acids typically hybridize under moderately stringent hybridization conditions. Exemplary "moderately stringent hybridization conditions" include a hybridization in a buffer of 40% formamide, 1 M NaCl, 1% SDS at 37°C, and a wash in 1X SSC at 45°C. A positive hybridization is at least twice background. Those of ordinary skill will readily recognize that alternative hybridization and wash conditions can be utilized to provide conditions of similar stringency. Additional guidelines for determining hybridization parameters are provided in numerous reference, e.g., and Current Protocols in Molecular Biology, ed. Ausubel, et al

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The phrase "functional effects" in the context of assays for testing compounds that modulate activity of an angiogenesis protein includes the determination of a parameter that is indirectly or directly under the influence of the angiogenesis protein, e.g., a functional, physical, or chemical effect, such as the ability to increase or decrease angiogenesis. It includes binding activity, the ability of cells to proliferate, expression in cells undergoing angiogenesis, and other characteristics of angiogenic cells. "Functional effects" include in vitro, in vivo, and ex vivo activities.

By "determining the functional effect" is meant assaying for a compound that increases or decreases a parameter that is indirectly or directly under the influence of an angiogenesis protein sequence, e.g., functional, physical and chemical effects. Such functional effects can be measured by any means known to those skilled in the art, e.g., changes in spectroscopic characteristics (e.g., fluorescence, absorbance, refractive index), hydrodynamic (e.g., shape), chromatographic, or solubility properties for the protein, measuring inducible markers or transcriptional activation of the angiogenesis protein; measuring binding activity or binding assays, e.g. binding to antibodies, and measuring cellular proliferation, particularly endothelial cell proliferation, cell viability, cell division especially of endothelial cells, lumen formation and capillary or vessel growth or formation. Determination of the functional effect of a compound on angiogenesis can also be performed using angiogenesis assays known to those of skill in the art such as an in vitro assays, e.g., in vitro endothelial cell tube formation assays, and other assays such as the chick CAM assay, the mouse corneal assay, and assays that assess vascularization of an implanted tumor. The

functional effects can be evaluated by many means known to those skilled in the art, e.g., microscopy for quantitative or qualitative measures of alterations in morphological features, e.g., tube or blood vessel formation, measurement of changes in RNA or protein levels for angiogenesis-associated sequences, measurement of RNA stability, identification of downstream or reporter gene expression (CAT, luciferase, β -gal, GFP and the like), e.g., via chemiluminescence, fluorescence, colorimetric reactions, antibody binding, inducible markers, and ligand binding assays.

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"Inhibitors", "activators", and "modulators" of angiogenic polynucleotide and polypeptide sequences are used to refer to activating, inhibitory, or modulating molecules identified using in vitro and in vivo assays of angiogenic polynucleotide and polypeptide sequences. Inhibitors are compounds that, e.g., bind to, partially or totally block activity, decrease, prevent, delay activation, inactivate, desensitize, or down regulate the activity or expression of angiogenesis proteins, e.g., antagonists. "Activators" are compounds that increase, open, activate, facilitate, enhance activation, sensitize, agonize, or up regulate angiogenesis protein activity. Inhibitors, activators, or modulators also include genetically modified versions of angiogenesis proteins, e.g., versions with altered activity, as well as naturally occurring and synthetic ligands, antagonists, agonists, antibodies, small chemical molecules and the like. Such assays for inhibitors and activators include, e.g., expressing the angiogenic protein in vitro, in cells, or cell membranes, applying putative modulator compounds, and then determining the functional effects on activity, as described above. Activators and inhibitors of angiogenesis can also be identified by incubating angiogenic cells with the test compound and determining increases or decreases in the expression of 1 or more angiogenesis proteins, e.g., 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 40, 50 or more angiogenesis proteins, such as angiogenesis proteins comprising the sequences set out in Table 8.

Samples or assays comprising angiogenesis proteins that are treated with a potential activator, inhibitor, or modulator are compared to control samples without the inhibitor, activator, or modulator to examine the extent of inhibition. Control samples (untreated with inhibitors) are assigned a relative protein activity value of 100%. Inhibition of a polypeptide is achieved when the activity value relative to the control is about 80%, preferably 50%, more preferably 25-0%. Activation of an angiogenesis polypeptide is achieved when the activity value relative to the control (untreated with activators) is 110%, more preferably 150%, more preferably 200-500% (i.e., two to five fold higher relative to the control), more preferably 1000-3000% higher.

"Antibody" refers to a polypeptide comprising a framework region from an immunoglobulin gene or fragments thereof that specifically binds and recognizes an antigen. The recognized immunoglobulin genes include the kappa, lambda, alpha, gamma, delta, epsilon, and mu constant region genes, as well as the myriad immunoglobulin variable region genes. Light chains are classified as either kappa or lambda. Heavy chains are classified as gamma, mu, alpha, delta, or epsilon, which in turn define the immunoglobulin classes, IgG, IgM, IgA, IgD and IgE, respectively. Typically, the antigen-binding region of an antibody will be most critical in specificity and affinity of binding.

An exemplary immunoglobulin (antibody) structural unit comprises a tetramer. Each tetramer is composed of two identical pairs of polypeptide chains, each pair having one "light" (about 25 kD) and one "heavy" chain (about 50-70 kD). The N-terminus of each chain defines a variable region of about 100 to 110 or more amino acids primarily responsible for antigen recognition. The terms variable light chain (V_L) and variable heavy chain (V_H) refer to these light and heavy chains respectively.

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Antibodies exist, e.g., as intact immunoglobulins or as a number of well-characterized fragments produced by digestion with various peptidases. Thus, for example, pepsin digests an antibody below the disulfide linkages in the hinge region to produce F(ab)'₂, a dimer of Fab which itself is a light chain joined to V_H-C_H1 by a disulfide bond. The F(ab)'₂ may be reduced under mild conditions to break the disulfide linkage in the hinge region, thereby converting the F(ab)'₂ dimer into an Fab' monomer. The Fab' monomer is essentially Fab with part of the hinge region (see Fundamental Immunology (Paul ed., 3d ed. 1993). While various antibody fragments are defined in terms of the digestion of an intact antibody, one of skill will appreciate that such fragments may be synthesized de novo either chemically or by using recombinant DNA methodology. Thus, the term antibody, as used herein, also includes antibody fragments either produced by the modification of whole antibodies, or those synthesized de novo using recombinant DNA methodologies (e.g., single chain Fv) or those identified using phage display libraries (see, e.g., McCafferty et al., Nature 348:552-554 (1990))

For preparation of antibodies, e.g., recombinant, monoclonal, or polyclonal antibodies, many technique known in the art can be used (see, e.g., Kohler & Milstein, Nature 256:495-497 (1975); Kozbor et al., Immunology Today 4: 72 (1983); Cole et al., pp. 77-96 in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc. (1985); Coligan, Current Protocols in Immunology (1991); Harlow & Lane, Antibodies, A Laboratory Manual (1988); and Goding, Monoclonal Antibodies: Principles and Practice (2d ed. 1986)).

Techniques for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce antibodies to polypeptides of this invention. Also, transgenic mice, or other organisms such as other mammals, may be used to express humanized antibodies. Alternatively, phage display technology can be used to identify antibodies and heteromeric Fab fragments that specifically bind to selected antigens (see, e.g., McCafferty et al., Nature 348:552-554 (1990); Marks et al., Biotechnology 10:779-783 (1992)).

A "chimeric antibody" is an antibody molecule in which (a) the constant region, or a portion thereof, is altered, replaced or exchanged so that the antigen binding site (variable region) is linked to a constant region of a different or altered class, effector function and/or species, or an entirely different molecule which confers new properties to the chimeric antibody, e.g., an enzyme, toxin, hormone, growth factor, drug, etc.; or (b) the variable region, or a portion thereof, is altered, replaced or exchanged with a variable region having a different or altered antigen specificity.

The detailed description of the invention includes discussion of the following

15 aspects of the invention:

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Expression of angiogenesis-associated sequences

Informatics

Angiogenesis-associated sequences

Detection of angiogenesis sequence for diagnostic and therapeutic applications

Modulators of angiogenesis

Methods of identifying variant angiogenesis-associated sequences

Administration of pharmaceutical and vaccine compositions Kits for use in diagnostic and/or prognostic applications.

25 Expression of angiogenesis-associated sequences

In one aspect, the expression levels of genes are determined in different patient samples for which diagnosis information is desired, to provide expression profiles. An expression profile of a particular sample is essentially a "fingerprint" of the state of the sample; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is unique to the state of the cell. That is, normal tissue may be distinguished from AD tissue. By comparing expression profiles of tissue in known different angiogenesis states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. The identification of sequences that are

differentially expressed in angiogenic versus non-angiogenic tissue allows the use of this information in a number of ways. For example, a particular treatment regime may be evaluated: does a chemotherapeutic drug act to down-regulate angiogenesis, and thus tumor growth or recurrence, in a particular patient. Similarly, diagnosis and treatment outcomes may be done or confirmed by comparing patient samples with the known expression profiles. Angiogenic tissue can also be analyzed to determine the stage of angiogenesis in the tissue. Furthermore, these gene expression profiles (or individual genes) allow screening of drug candidates with an eye to mimicking or altering a particular expression profile; for example, screening can be done for drugs that suppress the angiogenic expression profile. This may be done by making biochips comprising sets of the important angiogenesis genes, which can then be used in these screens. These methods can also be done on the protein basis; that is, protein expression levels of the angiogenic proteins can be evaluated for diagnostic purposes or to screen candidate agents. In addition, the angiogenic nucleic acid sequences can be administered for gene therapy purposes, including the administration of antisense nucleic acids, or the angiogenic proteins (including antibodies and other modulators thereof) administered as therapeutic drugs.

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Thus the present invention provides nucleic acid and protein sequences that are differentially expressed in angiogenesis, herein termed "angiogenesis sequences". As outlined below, angiogenesis sequences include those that are up-regulated (i.e. expressed at a higher level) in disorders associated with angiogenesis, as well as those that are down-regulated (i.e. expressed at a lower level). In a preferred embodiment, the angiogenesis sequences are from humans; however, as will be appreciated by those in the art, angiogenesis sequences from other organisms may be useful in animal models of disease and drug evaluation; thus, other angiogenesis sequences are provided, from vertebrates, including mammals, including rodents (rats, mice, hamsters, guinea pigs, etc.), primates, farm animals (including sheep, goats, pigs, cows, horses, etc). Angiogenesis sequences from other organisms may be obtained using the techniques outlined below.

Angiogenesis sequences can include both nucleic acid and amino acid sequences. In a preferred embodiment, the angiogenesis sequences are recombinant nucleic acids. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed in vitro, in general, by the manipulation of nucleic acid e.g., using polymerases and endonucleases, in a form not normally found in nature. Thus an isolated nucleic acid, in a linear form, or an expression vector formed in vitro by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this invention. It is

understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, *i.e.* using the *in vivo* cellular machinery of the host cell rather than *in vitro* manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention.

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Similarly, a "recombinant protein" is a protein made using recombinant techniques, i.e. through the expression of a recombinant nucleic acid as depicted above. A recombinant protein is distinguished from naturally occurring protein by at least one or more characteristics. For example, the protein may be isolated or purified away from some or all of the proteins and compounds with which it is normally associated in its wild type host, and thus may be substantially pure. For example, an isolated protein is unaccompanied by at least some of the material with which it is normally associated in its natural state, preferably constituting at least about 0.5%, more preferably at least about 5% by weight of the total protein in a given sample. A substantially pure protein comprises at least about 75% by weight of the total protein, with at least about 80% being preferred, and at least about 90% being particularly preferred. The definition includes the production of an angiogenesis protein from one organism in a different organism or host cell. Alternatively, the protein may be made at a significantly higher concentration than is normally seen, through the use of an inducible promoter or high expression promoter, such that the protein is made at increased concentration levels. Alternatively, the protein may be in a form not normally found in nature, as in the addition of an epitope tag or amino acid substitutions, insertions and deletions, as discussed below.

In a preferred embodiment, the angiogenesis sequences are nucleic acids. As will be appreciated by those in the art and is more fully outlined below, angiogenesis sequences are useful in a variety of applications, including diagnostic applications, which will detect naturally occurring nucleic acids, as well as screening applications; for example, biochips comprising nucleic acid probes to the angiogenesis sequences can be generated. In the broadest sense, then, by "nucleic acid" or "oligonucleotide" or grammatical equivalents herein means at least two nucleotides covalently linked together. A nucleic acid of the present invention will generally contain phosphodiester bonds, although in some cases, nucleic acid analogs are included that may have alternate backbones, comprising, for example, phosphoramidate, phosphorothioate, phosphorodithioate, or Omethylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press); and peptide nucleic acid backbones and linkages. Other

analog nucleic acids include those with positive backbones; non-ionic backbones, and non-ribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook. Nucleic acids containing one or more carbocyclic sugars are also included within one definition of nucleic acids. Modifications of the ribose-phosphate backbone may be done for a variety of reasons, for example to increase the stability and half-life of such molecules in physiological environments or as probes on a biochip.

As will be appreciated by those in the art, nucleic acid analogs may find use in the present invention. In addition, mixtures of naturally occurring nucleic acids and analogs can be made; alternatively, mixtures of different nucleic acid analogs, and mixtures of naturally occurring nucleic acids and analogs may be made.

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Particularly preferred are peptide nucleic acids (PNA) which includes peptide nucleic acid analogs. These backbones are substantially non-ionic under neutral conditions, in contrast to the highly charged phosphodiester backbone of naturally occurring nucleic acids. This results in two advantages. First, the PNA backbone exhibits improved hybridization kinetics. PNAs have larger changes in the melting temperature (Tm) for mismatched versus perfectly matched basepairs. DNA and RNA typically exhibit a 2-4°C drop in T_m for an internal mismatch. With the non-ionic PNA backbone, the drop is closer to 7-9°C. Similarly, due to their non-ionic nature, hybridization of the bases attached to these backbones is relatively insensitive to salt concentration. In addition, PNAs are not degraded by cellular enzymes, and thus can be more stable.

The nucleic acids may be single stranded or double stranded, as specified, or contain portions of both double stranded or single stranded sequence. As will be appreciated by those in the art, the depiction of a single strand also defines the sequence of the complementary strand; thus the sequences described herein also provide the complement of the sequence. The nucleic acid may be DNA, both genomic and cDNA, RNA or a hybrid, where the nucleic acid may contain combinations of deoxyribo- and ribo-nucleotides, and combinations of bases, including uracil, adenine, thymine, cytosine, guanine, inosine, xanthine hypoxanthine, isocytosine, isoguanine, etc. As used herein, the term "nucleoside" includes nucleotides and nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures. Thus for example the individual units of a peptide nucleic acid, each containing a base, are referred to herein as a nucleoside.

An angiogenesis sequence can be initially identified by substantial nucleic acid and/or amino acid sequence homology to the angiogenesis sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

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For identifying angiogenesis-associated sequences, the angiogenesis screen typically includes comparing genes identified in a modification of an *in vitro* model of angiogenesis as described in Hiraoka, Cell 95:365 (1998) with genes identified in controls. Samples of normal tissue and tissue undergoing angiogenesis are applied to biochips comprising nucleic acid probes. The samples are first microdissected, if applicable, and treated as is known in the art for the preparation of mRNA. Suitable biochips are commercially available, for example from Affymetrix. Gene expression profiles as described herein are generated and the data analyzed.

In a preferred embodiment, the genes showing changes in expression as

between normal and disease states are compared to genes expressed in other normal tissues, including, but not limited to lung, heart, brain, liver, breast, kidney, muscle, prostate, small intestine, large intestine, spleen, bone and placenta. In a preferred embodiment, those genes identified during the angiogenesis screen that are expressed in any significant amount in other tissues are removed from the profile, although in some embodiments, this is not necessary.

That is, when screening for drugs, it is usually preferable that the target be disease specific, to minimize possible side effects.

In a preferred embodiment, angiogenesis sequences are those that are upregulated in angiogenesis disorders; that is, the expression of these genes is higher in the
disease tissue as compared to normal tissue. "Up-regulation" as used herein means at least
about a two-fold change, preferably at least about a three fold change, with at least about
five-fold or higher being preferred. All accession numbers herein are for the GenBank
sequence database and the sequences of the accession numbers are hereby expressly
incorporated by reference. GenBank is known in the art, see, e.g., Benson, DA, et al.,
Nucleic Acids Research 26:1-7 (1998) and http://www.ncbi.nlm.nih.gov/. Sequences are also
avialable in other databases, e.g., European Molecular Biology Laboratory (EMBL) and
DNA Database of Japan (DDBJ). In addition, most preferred genes were found to be
expressed in a limited amount or not at all in heart, brain, lung, liver, breast, kidney, prostate,
small intestine and spleen.

In another preferred embodiment, angiogenesis sequences are those that are down-regulated in the angiogenesis disorder; that is, the expression of these genes is lower in angiogenic tissue as compared to normal tissue. "Down-regulation" as used herein means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred.

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Angiogenesis sequences according to the invention may be classified into discrete clusters of sequences based on common expression profiles of the sequences. Expression levels of angiogenesis sequences may increase or decrease as a function of time in a manner that correlates with the induction of angiogenesis. Alternatively, expression levels of angiogenesis sequences may both increase and decrease as a function of time. For example, expression levels of some angiogenesis sequences are temporarily induced or diminished during the switch to the angiogenesis phenotype, followed by a return to baseline expression levels. Tables 1-8 provides genes, the mRNA expression of which varies as a function of time in angiogenesis tissue when compared to normal tissue.

In a particularly preferred embodiment, angiogenesis sequences are those that are induced for a period of time, typically by positive angiogenic factors, followed by a return to the baseline levels. Sequences that are temporarily induced provide a means to target angiogenesis tissue, for example neovascularized tumors, at a particular stage of angiogenesis, while avoiding rapidly growing tissue that require perpetual vascularization. Such positive angiogenic factors include α FGF, β FGF, VEGF, angiogenin and the like.

Induced angiogenesis sequences also are further categorized with respect to the timing of induction. For example, some angiogenesis genes may be induced at an early time period, such as within 10 minutes of the induction of angiogenesis. Others may be induced later, such as between 5 and 60 minutes, while yet others may be induced for a time period of about two hours or more followed by a return to baseline expression levels.

In another preferred embodiment are angiogenesis sequences that are inhibited or reduced as a function of time followed by a return to "normal" expression levels.

Inhibitors of angiogenesis are examples of molecules that have this expression profile. These sequences also can be further divided into groups depending on the timing of diminished expression. For example, some molecules may display reduced expression within 10 minutes of the induction of angiogenesis. Others may be diminished later, such as between 5 and 60 minutes, while others may be diminished for a time period of about two hours or more

followed by a return to baseline. Examples of such negative angiogenic factors include thrombospondin and endostatin to name a few.

In yet another preferred embodiment are angiogenesis sequences that are induced for prolonged periods. These sequences are typically associated with induction of angiogenesis and may participate in induction and/or maintenance of the angiogenesis phenotype.

In another preferred embodiment are angiogenesis sequences, the expression of which is reduced or diminished for prolonged periods in angiogenic tissue. These sequences are typically angiogenesis inhibitors and their diminution is correlated with an increase in angiogenesis.

Informatics

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The ability to identify genes that undergo changes in expression with time during angiogenesis can additionally provide high-resolution, high-sensitivity datasets which can be used in the areas of diagnostics, therapeutics, drug development, biosensor development, and other related areas. For example, the expression profiles can be used in diagnostic or prognostic evaluation of patients with angiogenesis-associated disease. Or as another example, subcellular toxicological information can be generated to better direct drug structure and activity correlation (see, Anderson, L., "Pharmaceutical Proteomics: Targets, Mechanism, and Function," paper presented at the IBC Proteomics conference, Coronado, CA (June 11-12, 1998)). Subcellular toxicological information can also be utilized in a biological sensor device to predict the likely toxicological effect of chemical exposures and likely tolerable exposure thresholds (see, U.S. Patent No. 5,811,231). Similar advantages accrue from datasets relevant to other biomolecules and bioactive agents (e.g., nucleic acids, saccharides, lipids, drugs, and the like).

Thus, in another embodiment, the present invention provides a database that includes at least one set of data assay data. The data contained in the database is acquired, e.g., using array analysis either singly or in a library format. The database can be in substantially any form in which data can be maintained and transmitted, but is preferably an electronic database. The electronic database of the invention can be maintained on any electronic device allowing for the storage of and access to the database, such as a personal computer, but is preferably distributed on a wide area network, such as the World Wide Web.

The focus of the present section on databases that include peptide sequence data is for clarity of illustration only. It will be apparent to those of skill in the art that similar databases can be assembled for any assay data acquired using an assay of the invention.

The compositions and methods for identifying and/or quantitating the relative and/or absolute abundance of a variety of molecular and macromolecular species from a biological sample undergoing angiogenesis, *i.e.*, the identification of angiogenesis-associated sequences described herein, provide an abundance of information, which can be correlated with pathological conditions, predisposition to disease, drug testing, therapeutic monitoring, gene-disease causal linkages, identification of correlates of immunity and physiological status, among others. Although the data generated from the assays of the invention is suited for manual review and analysis, in a preferred embodiment, prior data processing using high-speed computers is utilized.

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An array of methods for indexing and retrieving biomolecular information is known in the art. For example, U.S. Patents 6,023,659 and 5,966,712 disclose a relational database system for storing biomolecular sequence information in a manner that allows sequences to be catalogued and searched according to one or more protein function hierarchies. U.S. Patent 5,953,727 discloses a relational database having sequence records containing information in a format that allows a collection of partial-length DNA sequences to be catalogued and searched according to association with one or more sequencing projects for obtaining full-length sequences from the collection of partial length sequences. U.S. Patent 5,706,498 discloses a gene database retrieval system for making a retrieval of a gene sequence similar to a sequence data item in a gene database based on the degree of similarity between a key sequence and a target sequence. U.S. Patent 5,538,897 discloses a method using mass spectroscopy fragmentation patterns of peptides to identify amino acid sequences in computer databases by comparison of predicted mass spectra with experimentally-derived mass spectra using a closeness-of-fit measure. U.S. Patent 5,926,818 discloses a multidimensional database comprising a functionality for multi-dimensional data analysis described as on-line analytical processing (OLAP), which entails the consolidation of projected and actual data according to more than one consolidation path or dimension. U.S. Patent 5,295,261 reports a hybrid database structure in which the fields of each database record are divided into two classes, navigational and informational data, with navigational fields stored in a hierarchical topological map which can be viewed as a tree structure or as the merger of two or more such tree structures.

The present invention provides a computer database comprising a computer and software for storing in computer-retrievable form assay data records cross-tabulated, e.g., with data specifying the source of the target-containing sample from which each sequence specificity record was obtained.

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In an exemplary embodiment, at least one of the sources of target-containing sample is from a control tissue sample known to be free of pathological disorders. In a variation, at least one of the sources is a known pathological tissue specimen, e.g., a neoplastic lesion or another tissue specimen to be analyzed for angiogenesis. In another variation, the assay records cross-tabulate one or more of the following parameters for each target species in a sample: (1) a unique identification code, which can include, e.g., a target molecular structure and/or characteristic separation coordinate (e.g., electrophoretic coordinates); (2) sample source; and (3) absolute and/or relative quantity of the target species present in the sample.

The invention also provides for the storage and retrieval of a collection of target data in a computer data storage apparatus, which can include magnetic disks, optical disks, magneto-optical disks, DRAM, SRAM, SGRAM, SDRAM, RDRAM, DDR RAM, magnetic bubble memory devices, and other data storage devices, including CPU registers and on-CPU data storage arrays. Typically, the target data records are stored as a bit pattern in an array of magnetic domains on a magnetizable medium or as an array of charge states or transistor gate states, such as an array of cells in a DRAM device (e.g., each cell comprised of a transistor and a charge storage area, which may be on the transistor). In one embodiment, the invention provides such storage devices, and computer systems built therewith, comprising a bit pattern encoding a protein expression fingerprint record comprising unique identifiers for at least 10 target data records cross-tabulated with target source.

When the target is a peptide or nucleic acid, the invention preferably provides a method for identifying related peptide or nucleic acid sequences, comprising performing a computerized comparison between a peptide or nucleic acid sequence assay record stored in or retrieved from a computer storage device or database and at least one other sequence. The comparison can include a sequence analysis or comparison algorithm or computer program embodiment thereof (e.g., FASTA, TFASTA, GAP, BESTFIT) and/or the comparison may be of the relative amount of a peptide or nucleic acid sequence in a pool of sequences determined from a polypeptide or nucleic acid sample of a specimen.

The invention also preferably provides a magnetic disk, such as an IBM-compatible (DOS, Windows, Windows95/98/2000, Windows NT, OS/2) or other format

(e.g., Linux, SunOS, Solaris, AIX, SCO Unix, VMS, MV, Macintosh, etc.) floppy diskette or hard (fixed, Winchester) disk drive, comprising a bit pattern encoding data from an assay of the invention in a file format suitable for retrieval and processing in a computerized sequence analysis, comparison, or relative quantitation method.

The invention also provides a network, comprising a plurality of computing devices linked via a data link, such as an Ethernet cable (coax or 10BaseT), telephone line, ISDN line, wireless network, optical fiber, or other suitable signal tranmission medium, whereby at least one network device (e.g., computer, disk array, etc.) comprises a pattern of magnetic domains (e.g., magnetic disk) and/or charge domains (e.g., an array of DRAM cells) composing a bit pattern encoding data acquired from an assay of the invention.

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The invention also provides a method for transmitting assay data that includes generating an electronic signal on an electronic communications device, such as a modem, ISDN terminal adapter, DSL, cable modem, ATM switch, or the like, wherein the signal includes (in native or encrypted format) a bit pattern encoding data from an assay or a database comprising a plurality of assay results obtained by the method of the invention.

In a preferred embodiment, the invention provides a computer system for comparing a query target to a database containing an array of data structures, such as an assay result obtained by the method of the invention, and ranking database targets based on the degree of identity and gap weight to the target data. A central processor is preferably initialized to load and execute the computer program for alignment and/or comparison of the assay results. Data for a query target is entered into the central processor via an I/O device. Execution of the computer program results in the central processor retrieving the assay data from the data file, which comprises a binary description of an assay result.

The target data or record and the computer program can be transferred to secondary memory, which is typically random access memory (e.g., DRAM, SRAM, SGRAM, or SDRAM). Targets are ranked according to the degree of correspondence between a selected assay characteristic (e.g., binding to a selected affinity moiety) and the same characteristic of the query target and results are output via an I/O device. For example, a central processor can be a conventional computer (e.g., Intel Pentium, PowerPC, Alpha, PA-8000, SPARC, MIPS 4400, MIPS 10000, VAX, etc.); a program can be a commercial or public domain molecular biology software package (e.g., UWGCG Sequence Analysis Software, Darwin); a data file can be an optical or magnetic disk, a data server, a memory device (e.g., DRAM, SRAM, SGRAM, SDRAM, EPROM, bubble memory, flash memory, etc.); an I/O device can be a terminal comprising a video display and a keyboard, a modem,

an ISDN terminal adapter, an Ethernet port, a punched card reader, a magnetic strip reader, or other suitable I/O device.

The invention also preferably provides the use of a computer system, such as that described above, which comprises: (1) a computer; (2) a stored bit pattern encoding a collection of peptide sequence specificity records obtained by the methods of the invention, which may be stored in the computer; (3) a comparison target, such as a query target; and (4) a program for alignment and comparison, typically with rank-ordering of comparison results on the basis of computed similarity values.

10 Angiogenesis-associated sequences

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Angiogenesis proteins of the present invention may be classified as secreted proteins, transmembrane proteins or intracellular proteins. In one embodiment, the angiogenesis protein is an intracellular protein. Intracellular proteins may be found in the cytoplasm and/or in the nucleus or associated with the intracellular side of the plasma membrane. Intracellular proteins are involved in all aspects of cellular function and replication (including, e.g., signaling pathways); aberrant expression of such proteins often results in unregulated or disregulated cellular processes (see, e.g., Molecular Biology of the Cell, 3rd Edition, Alberts, Ed., Garland Pub., 1994). For example, many intracellular proteins have enzymatic activity such as protein kinase activity, protein phosphatase activity, protease activity, nucleotide cyclase activity, polymerase activity and the like. Intracellular proteins also serve as docking proteins that are involved in organizing complexes of proteins, or targeting proteins to various subcellular localizations, and are involved in maintaining the structural integrity of organelles.

An increasingly appreciated concept in characterizing proteins is the presence in the proteins of one or more motifs for which defined functions have been attributed. In addition to the highly conserved sequences found in the enzymatic domain of proteins, highly conserved sequences have been identified in proteins that are involved in protein-protein interaction. For example, Src-homology-2 (SH2) domains bind tyrosine-phosphorylated targets in a sequence dependent manner. PTB domains, which are distinct from SH2 domains, also bind tyrosine phosphorylated targets. SH3 domains bind to proline-rich targets. In addition, PH domains, tetratricopeptide repeats and WD domains to name only a few, have been shown to mediate protein-protein interactions. Some of these may also be involved in binding to phospholipids or other second messengers. As will be appreciated by one of ordinary skill in the art, these motifs can be identified on the basis of primary

sequence; thus, an analysis of the sequence of proteins may provide insight into both the enzymatic potential of the molecule and/or molecules with which the protein may associate.

In another embodiment, the angiogenesis sequences are transmembrane proteins. Transmembrane proteins are molecules that span a phospholipid bilayer of a cell. They may have an intracellular domain, an extracellular domain, or both. The intracellular domains of such proteins may have a number of functions including those already described for intracellular proteins. For example, the intracellular domain may have enzymatic activity and/or may serve as a binding site for additional proteins. Frequently the intracellular domain of transmembrane proteins serves both roles. For example certain receptor tyrosine kinases have both protein kinase activity and SH2 domains. In addition, autophosphorylation of tyrosines on the receptor molecule itself, creates binding sites for additional SH2 domain containing proteins.

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Transmembrane proteins may contain from one to many transmembrane domains. For example, receptor tyrosine kinases, certain cytokine receptors, receptor guanylyl cyclases and receptor serine/threonine protein kinases contain a single transmembrane domain. However, various other proteins including channels and adenylyl cyclases contain numerous transmembrane domains. Many important cell surface receptors such as G protein coupled receptors (GPCRs) are classified as "seven transmembrane domain" proteins, as they contain 7 membrane spanning regions. Characteristics of transmembrane domains include approximately 20 consecutive hydrophobic amino acids that may be followed or flanked by charged amino acids. Therefore, upon analysis of the amino acid sequence of a particular protein, the localization and number of transmembrane domains within the protein may be predicted (see, e.g. PSORT web site http://psort.nibb.ac.jp/).

The extracellular domains of transmembrane proteins are diverse; however, conserved motifs are found repeatedly among various extracellular domains. Conserved structure and/or functions have been ascribed to different extracellular motifs. Many extracellular domains are involved in binding to other molecules. In one aspect, extracellular domains are found on receptors. Factors that bind the receptor domain include circulating ligands, which may be peptides, proteins, or small molecules such as adenosine and the like. For example, growth factors such as EGF, FGF and PDGF are circulating growth factors that bind to their cognate receptors to initiate a variety of cellular responses. Other factors include cytokines, mitogenic factors, neurotrophic factors and the like. Extracellular domains also bind to cell-associated molecules. In this respect, they mediate cell-cell interactions. Cell-associated ligands can be tethered to the cell for example via a glycosylphosphatidylinositol

(GPI) anchor, or may themselves be transmembrane proteins. Extracellular domains also associate with the extracellular matrix and contribute to the maintenance of the cell structure.

Angiogenesis proteins that are transmembrane are particularly preferred in the present invention as they are readily accessible targets for immunotherapeutics, as are described herein. In addition, as outlined below, transmembrane proteins can be also useful in imaging modalities. Antibodies may be used to label such readily accessible proteins in situ. Alternatively, antibodies can also label intracellular proteins, in which case samples are typically permeablized to provide acess to intracellular proteins.

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It will also be appreciated by those in the art that a transmembrane protein can be made soluble by removing transmembrane sequences, for example through recombinant methods. Furthermore, transmembrane proteins that have been made soluble can be made to be secreted through recombinant means by adding an appropriate signal sequence.

In another embodiment, the angiogenesis proteins are secreted proteins; the secretion of which can be either constitutive or regulated. These proteins have a signal peptide or signal sequence that targets the molecule to the secretory pathway. Secreted proteins are involved in numerous physiological events; by virtue of their circulating nature, they serve to transmit signals to various other cell types. The secreted protein may function in an autocrine manner (acting on the cell that secreted the factor), a paracrine manner (acting on cells in close proximity to the cell that secreted the factor) or an endocrine manner (acting on cells at a distance). Thus secreted molecules find use in modulating or altering numerous aspects of physiology. Angiogenesis proteins that are secreted proteins are particularly preferred in the present invention as they serve as good targets for diagnostic markers, e.g., for blood or serum tests.

An angiogenesis sequence is typically initially identified by substantial nucleic acid and/or amino acid sequence homology or linkage to the angiogenesis sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions. Typically, linked sequences on a mRNA are found on the same molecule.

As detailed in the definitions, percent identity can be determined using an algorithm such as BLAST. A preferred method utilizes the BLASTN module of WU-BLAST-2 set to the default parameters, with overlap span and overlap fraction set to 1 and 0.125, respectively. The alignment may include the introduction of gaps in the sequences to be aligned. In addition, for sequences which contain either more or fewer nucleotides than

those of the nucleic acids of the figures, it is understood that the percentage of homology will be determined based on the number of homologous nucleosides in relation to the total number of nucleosides. Thus, for example, homology of sequences shorter than those of the sequences identified herein and as discussed below, will be determined using the number of nucleosides in the shorter sequence.

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In one embodiment, the nucleic acid homology is determined through hybridization studies. Thus, e.g., nucleic acids which hybridize under high stringency to a nucleic acid of Tables 1-8, or its complement, or is also found on naturally occurring mRNAs is considered an angiogenesis sequence. In another embodiment, less stringent hybridization conditions are used; for example, moderate or low stringency conditions may be used, as are known in the art; see Ausubel, supra, and Tijssen, supra.

In addition, the angiogenesis nucleic acid sequences of the invention, e.g, the sequence in Tables 1-8, are fragments of larger genes, *i.e.* they are nucleic acid segments. "Genes" in this context includes coding regions, non-coding regions, and mixtures of coding and non-coding regions. Accordingly, as will be appreciated by those in the art, using the sequences provided herein, extended sequences, in either direction, of the angiogenesis genes can be obtained, using techniques well known in the art for cloning either longer sequences or the full length sequences; see Ausubel, *et al.*, *supra*. Much can be done by informatics and many sequences can be clustered to include multiple sequences, *e.g.*, systems such as UniGene (see, http://www.ncbi.nlm.nih.gov/UniGene/).

Once the angiogenesis nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire angiogenesis nucleic acid coding regions or the entire mRNA sequence. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant angiogenesis nucleic acid can be further-used as a probe to identify and isolate other angiogenesis nucleic acids, for example extended coding regions. It can also be used as a "precursor" nucleic acid to make modified or variant angiogenesis nucleic acids and proteins.

The angiogenesis nucleic acids of the present invention are used in several ways. In a first embodiment, nucleic acid probes to the angiogenesis nucleic acids are made and attached to biochips to be used in screening and diagnostic methods, as outlined below, or for administration, for example for gene therapy, vaccine, and/or antisense applications. Alternatively, the angiogenesis nucleic acids that include coding regions of angiogenesis

proteins can be put into expression vectors for the expression of angiogenesis proteins, again for screening purposes or for administration to a patient.

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In a preferred embodiment, nucleic acid probes to angiogenesis nucleic acids (both the nucleic acid sequences outlined in the figures and/or the complements thereof) are made. The nucleic acid probes attached to the biochip are designed to be substantially complementary to the angiogenesis nucleic acids, *i.e.* the target sequence (either the target sequence of the sample or to other probe sequences, for example in sandwich assays), such that hybridization of the target sequence and the probes of the present invention occurs. As outlined below, this complementarity need not be perfect; there may be any number of base pair mismatches which will interfere with hybridization between the target sequence and the single stranded nucleic acids of the present invention. However, if the number of mutations is so great that no hybridization can occur under even the least stringent of hybridization conditions, the sequence is not a complementary target sequence. Thus, by "substantially complementary" herein is meant that the probes are sufficiently complementary to the target sequences to hybridize under normal reaction conditions, particularly high stringency conditions, as outlined herein.

A nucleic acid probe is generally single stranded but can be partially single and partially double stranded. The strandedness of the probe is dictated by the structure, composition, and properties of the target sequence. In general, the nucleic acid probes range from about 8 to about 100 bases long, with from about 10 to about 80 bases being preferred, and from about 30 to about 50 bases being particularly preferred. That is, generally whole genes are not used. In some embodiments, much longer nucleic acids can be used, up to hundreds of bases.

In a preferred embodiment, more than one probe per sequence is used, with either overlapping probes or probes to different sections of the target being used. That is, two, three, four or more probes, with three being preferred, are used to build in a redundancy for a particular target. The probes can be overlapping (i.e. have some sequence in common), or separate. In some cases, PCR primers may be used to amplify signal for higher sensitivity.

As will be appreciated by those in the art, nucleic acids can be attached or immobilized to a solid support in a wide variety of ways. By "immobilized" and grammatical equivalents herein is meant the association or binding between the nucleic acid probe and the solid support is sufficient to be stable under the conditions of binding, washing, analysis, and removal as outlined below. The binding can typically be covalent or non-covalent. By "non-covalent binding" and grammatical equivalents herein is meant one or more of electrostatic,

hydrophilic, and hydrophobic interactions. Included in non-covalent binding is the covalent attachment of a molecule, such as, streptavidin to the support and the non-covalent binding of the biotinylated probe to the streptavidin. By "covalent binding" and grammatical equivalents herein is meant that the two moieties, the solid support and the probe, are attached by at least one bond, including sigma bonds, pi bonds and coordination bonds. Covalent bonds can be formed directly between the probe and the solid support or can be formed by a cross linker or by inclusion of a specific reactive group on either the solid support or the probe or both molecules. Immobilization may also involve a combination of covalent and non-covalent interactions.

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In general, the probes are attached to the biochip in a wide variety of ways, as will be appreciated by those in the art. As described herein, the nucleic acids can either be synthesized first, with subsequent attachment to the biochip, or can be directly synthesized on the biochip.

The biochip comprises a suitable solid substrate. By "substrate" or "solid support" or other grammatical equivalents herein is meant a material that can be modified to contain discrete individual sites appropriate for the attachment or association of the nucleic acid probes and is amenable to at least one detection method. As will be appreciated by those in the art, the number of possible substrates are very large, and include, but are not limited to, glass and modified or functionalized glass, plastics (including acrylics, polystyrene and copolymers of styrene and other materials, polypropylene, polyethylene, polybutylene, polyurethanes, TeflonJ, etc.), polysaccharides, nylon or nitrocellulose, resins, silica or silicabased materials including silicon and modified silicon, carbon, metals, inorganic glasses, plastics, etc. In general, the substrates allow optical detection and do not appreciably fluorescese. A preferred substrate is described in copending application entitled Reusable Low Fluorescent Plastic Biochip, U.S. Application Serial No. 09/270,214, filed March 15, 1999, herein incorporated by reference in its entirety.

Generally the substrate is planar, although as will be appreciated by those in the art, other configurations of substrates may be used as well. For example, the probes may be placed on the inside surface of a tube, for flow-through sample analysis to minimize sample volume. Similarly, the substrate may be flexible, such as a flexible foam, including closed cell foams made of particular plastics.

In a preferred embodiment, the surface of the biochip and the probe may be derivatized with chemical functional groups for subsequent attachment of the two. Thus, for example, the biochip is derivatized with a chemical functional group including, but not

limited to, amino groups, carboxy groups, oxo groups and thiol groups, with amino groups being particularly preferred. Using these functional groups, the probes can be attached using functional groups on the probes. For example, nucleic acids containing amino groups can be attached to surfaces comprising amino groups, for example using linkers as are known in the art; for example, homo-or hetero-bifunctional linkers as are well known (see 1994 Pierce Chemical Company catalog, technical section on cross-linkers, pages 155-200, incorporated herein by reference). In addition, in some cases, additional linkers, such as alkyl groups (including substituted and heteroalkyl groups) may be used.

In this embodiment, oligonucleotides are synthesized as is known in the art, and then attached to the surface of the solid support. As will be appreciated by those skilled in the art, either the 5' or 3' terminus may be attached to the solid support, or attachment may be via an internal nucleoside.

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In another embodiment, the immobilization to the solid support may be very strong, yet non-covalent. For example, biotinylated oligonucleotides can be made, which bind to surfaces covalently coated with streptavidin, resulting in attachment.

Alternatively, the oligonucleotides may be synthesized on the surface, as is known in the art. For example, photoactivation techniques utilizing photopolymerization compounds and techniques are used. In a preferred embodiment, the nucleic acids can be synthesized in situ, using well known photolithographic techniques, such as those described in WO 95/25116; WO 95/35505; U.S. Patent Nos. 5,700,637 and 5,445,934; and references cited within, all of which are expressly incorporated by reference; these methods of attachment form the basis of the Affimetrix GeneChipTM technology.

Often, amplification-based assays are performed to measure the expression level of angiogenesis-associated sequences. These assays are typically performed in conjunction with reverse transcription. In such assays, an angiogenesis-associated nucleic acid sequence acts as a template in an amplification reaction (e.g., Polymerase Chain Reaction, or PCR). In a quantitative amplification, the amount of amplification product will be proportional to the amount of template in the original sample. Comparison to appropriate controls provides a measure of the amount of angiogenesis-associated RNA. Methods of quantitative amplification are well known to those of skill in the art. Detailed protocols for quantitative PCR are provided, e.g., in Innis et al. (1990) PCR Protocols, A Guide to Methods and Applications, Academic Press, Inc. N.Y.).

In some embodiments, a TaqMan based assay is used to measure expression. TaqMan based assays use a fluorogenic oligonucleotide probe that contains a 5' fluorescent

dye and a 3' quenching agent. The probe hybridizes to a PCR product, but cannot itself be extended due to a blocking agent at the 3' end. When the PCR product is amplified in subsequent cycles, the 5' nuclease activity of the polymerase, e.g., AmpliTaq, results in the cleavage of the TaqMan probe. This cleavage separates the 5' fluorescent dye and the 3' quenching agent, thereby resulting in an increase in fluorescence as a function of amplification (see, for example, literature provided by Perkin-Elmer, e.g., www2.perkin-elmer.com).

Other suitable amplification methods include, but are not limited to, ligase chain reaction (LCR) (see, Wu and Wallace (1989) Genomics 4: 560, Landegren et al. (1988) Science 241: 1077, and Barringer et al. (1990) Gene 89: 117), transcription amplification (Kwoh et al. (1989) Proc. Natl. Acad. Sci. USA 86: 1173), self-sustained sequence replication (Guatelli et al. (1990) Proc. Nat. Acad. Sci. USA 87: 1874), dot PCR, and linker adapter PCR, etc.

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In a preferred embodiment, angiogenesis nucleic acids, e.g., encoding angiogenesis proteins are used to make a variety of expression vectors to express angiogenesis proteins which can then be used in screening assays, as described below. Expression vectors and recombinant DNA technology are well known to those of skill in the art (see, e.g., Ausubel, supra, and Gene Expression Systems, Fernandez & Hoeffler, Eds, Academic Press, 1999) and are used to express proteins. The expression vectors may be either self-replicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these expression vectors include transcriptional and translational regulatory nucleic acid operably linked to the nucleic acid encoding the angiogenesis protein. The term "control sequences" refers to DNA sequences used for the expression of an operably linked coding sequence in a particular host organism. Control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation.

Generally, "operably linked" means that the DNA sequences being linked are contiguous,

and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is typically accomplished by ligation at convenient restriction sites. If such sites do not exist, synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. Transcriptional and translational regulatory nucleic acid will generally be appropriate to the host cell used to express the angiogenesis protein; for example, transcriptional and translational regulatory nucleic acid sequences from Bacillus are preferably used to express the angiogenesis protein in Bacillus. Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

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In general, transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, an expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, for example in mammalian or insect cells for expression and in a procaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating vector may be directed to a specific locus in the host cell by selecting the appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art (e.g., Fernandez & Hoeffler, supra). See also Kitamura, et al. (1995) PNAS 92:9146-9150.

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

The angiogenesis proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding an angiogenesis protein, under the appropriate conditions to induce or cause expression of the

angiogenesis protein. Conditions appropriate for angiogenesis protein expression will vary with the choice of the expression vector and the host cell, and will be easily ascertained by one skilled in the art through routine experimentation or optimization. For example, the use of constitutive promoters in the expression vector will require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions for induction. In addition, in some embodiments, the timing of the harvest is important. For example, the baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

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Appropriate host cells include yeast, bacteria, archaebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are Saccharomyces cerevisiae and other yeasts, E. coli, Bacillus subtilis, Sf9 cells, C129 cells, 293 cells, Neurospora, BHK, CHO, COS, HeLa cells, HUVEC (human umbilical vein endothelial cells), THP1 cells (a macrophage cell line) and various other human cells and cell lines.

In a preferred embodiment, the angiogenesis proteins are expressed in mammalian cells. Mammalian expression systems are also known in the art, and include retroviral and adenoviral systems. Of particular use as mammalian promoters are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, herpes simplex virus promoter, and the CMV promoter (see, e.g., Fernandez & Hoeffler, supra). Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. Examples of transcription terminator and polyadenlytion signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used.

Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, viral infection, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

In a preferred embodiment, angiogenesis proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art. Promoters from bacteriophage may also be used and are known in the art. In addition, synthetic promoters

and hybrid promoters are also useful; for example, the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. In addition to a functioning promoter sequence, an efficient ribosome binding site is desirable. The expression vector may also include a signal peptide sequence that provides for secretion of the angiogenesis protein in bacteria. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways. These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for Bacillus subtilis. E. coli, Streptococcus cremoris, and Streptococcus lividans, among others (e.g., Fernandez & Hoeffler, supra). The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

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In one embodiment, angiogenesis proteins are produced in insect cells.

Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art.

In a preferred embodiment, angiogenesis protein is produced in yeast cells. Yeast expression systems are well known in the art, and include expression vectors for Saccharomyces cerevisiae, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fragilis and K. lactis, Pichia guillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica.

The angiogenesis protein may also be made as a fusion protein, using techniques well known in the art. Thus, for example, for the creation of monoclonal antibodies, if the desired epitope is small, the angiogenesis protein may be fused to a carrier protein to form an immunogen. Alternatively, the angiogenesis protein may be made as a fusion protein to increase expression, or for other reasons. For example, when the angiogenesis protein is an angiogenesis peptide, the nucleic acid encoding the peptide may be linked to another nucleic acid for expression purposes. Fusion with detection epitope tags can be made, e.g., with FLAG, His 6, myc, HA, etc.

In one embodiment, the angiogenesis nucleic acids, proteins and antibodies of the invention are labeled. By "labeled" herein is meant that a compound has at least one element, isotope or chemical compound attached to enable the detection of the compound. In general, labels fall into three classes: a) isotopic labels, which may be radioactive or heavy isotopes; b) immune labels, which may be antibodies, antigens, or epitope tags and c) colored or fluorescent dyes. The labels may be incorporated into the angiogenesis nucleic acids, proteins and antibodies at any position. For example, the label should be capable of producing, either directly or indirectly, a detectable signal. The detectable moiety may be a radioisotope, such as ³H, ¹⁴C, ³²P, ³⁵S, or ¹²⁵I, a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin, or an enzyme, such as alkaline phosphatase, beta-galactosidase or horseradish peroxidase. Any method known in the art for conjugating the antibody to the label may be employed, including those methods described by Hunter et al., Nature, 144:945 (1962); David et al., Biochemistry, 13:1014 (1974); Pain et al., J. Immunol. Meth., 40:219 (1981); and Nygren, J. Histochem. and Cytochem., 30:407 (1982).

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Accordingly, the present invention also provides angiogenesis protein sequences. An angiogenesis protein of the present invention may be identified in several ways. "Protein" in this sense includes proteins, polypeptides, and peptides. As will be appreciated by those in the art, the nucleic acid sequences of the invention can be used to generate protein sequences. There are a variety of ways to do this, including cloning the entire gene and verifying its frame and amino acid sequence, or by comparing it to known sequences to search for homology to provide a frame, assuming the angiogenesis protein has an identifiable motif or homology to some protein in the database being used. Generally, the nucleic acid sequences are input into a program that will search all three frames for homology. This is done in a preferred embodiment using the following NCBI Advanced BLAST parameters. The program is blastx or blastn. The database is nr. The input data is as "Sequence in FASTA format". The organism list is "none". The "expect" is 10; the filter is default. The "descriptions" is 500, the "alignments" is 500, and the "alignment view" is pairwise. The "Query Genetic Codes" is standard (1). The matrix is BLOSUM62; gap existence cost is 11, per residue gap cost is 1; and the lambda ratio is .85 default. This results in the generation of a putative protein sequence.

Also included within one embodiment of angiogenesis proteins are amino acid variants of the naturally occurring sequences, as determined herein. Preferably, the variants are preferably greater than about 75% homologous to the wild-type sequence, more

preferably greater than about 80%, even more preferably greater than about 85% and most preferably greater than 90%. In some embodiments the homology will be as high as about 93 to 95 or 98%. As for nucleic acids, homology in this context means sequence similarity or identity, with identity being preferred. This homology will be determined using standard techniques well known in the art as are outlined above for the nucleic acid homologies.

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Angiogenesis proteins of the present invention may be shorter or longer than the wild type amino acid sequences. Thus, in a preferred embodiment, included within the definition of angiogenesis proteins are portions or fragments of the wild type sequences. herein. In addition, as outlined above, the angiogenesis nucleic acids of the invention may be used to obtain additional coding regions, and thus additional protein sequence, using techniques known in the art.

In a preferred embodiment, the angiogenesis proteins are derivative or variant angiogenesis proteins as compared to the wild-type sequence. That is, as outlined more fully below, the derivative angiogenesis peptide will often contain at least one amino acid substitution, deletion or insertion, with amino acid substitutions being particularly preferred. The amino acid substitution, insertion or deletion may occur at any residue within the angiogenesis peptide.

Also included within one embodiment of angiogenesis proteins of the present invention are amino acid sequence variants. These variants typically fall into one or more of three classes: substitutional, insertional or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the angiogenesis protein, using cassette or PCR mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture as outlined above. However, variant angiogenesis protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the angiogenesis protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed angiogenesis variants screened for

the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, for example, M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of angiogenesis protein activities.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger insertions may be tolerated. Deletions range from about 1 to about 20 residues, although in some cases deletions may be much larger.

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Substitutions, deletions, insertions or any combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the molecule. However, larger changes may be tolerated in certain circumstances. When small alterations in the characteristics of the angiogenesis protein are desired, substitutions are generally made in accordance with the amino acid substitution chart provided in the definition section.

Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those provided in the definition of "conservative substitution". For example, substitutions may be made which more significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g. seryl or threonyl, is substituted for (or by) a hydrophobic residue, e.g. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g. lysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g. glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g. phenylalanine, is substituted for (or by) one not having a side chain, e.g. glycine.

The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analog, although variants also are selected to modify the characteristics of the angiogenesis proteins as needed. Alternatively, the variant may be designed such that the biological activity of the angiogenesis protein is altered. For example, glycosylation sites may be altered or removed.

Covalent modifications of angiogenesis polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino

acid residues of an angiogenesis polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C-terminal residues of an angiogenesis polypeptide. Derivatization with bifunctional agents is useful, for instance, for crosslinking angiogenesis polypeptides to a water-insoluble support matrix or surface for use in the method for purifying anti-angiogenesis polypeptide antibodies or screening assays, as is more fully described below. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidylpropionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate.

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Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl, threonyl or tyrosyl residues, methylation of the γ -amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, Proteins: Structure and Molecular Properties, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the angiogenesis polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence angiogenesis polypeptide, and/or adding one or more glycosylation sites that are not present in the native sequence angiogenesis polypeptide. Glycosylation patterns can be altered in many ways. For example the use of different cell types to express angiogenesis-associated sequences can result in different glycosylation patterns.

Addition of glycosylation sites to angiogenesis polypeptides may also be accomplished by altering the amino acid sequence thereof. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence angiogenesis polypeptide (for O-linked glycosylation sites). The angiogenesis amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the angiogenesis polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the angiogenesis polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, CRC Crit. Rev. Biochem., pp. 259-306 (1981).

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Removal of carbohydrate moieties present on the angiogenesis polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo-and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

Another type of covalent modification of angiogenesis comprises linking the angiogenesis polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

Angiogenesis polypeptides of the present invention may also be modified in a way to form chimeric molecules comprising an angiogenesis polypeptide fused to another, heterologous polypeptide or amino acid sequence. In one embodiment, such a chimeric molecule comprises a fusion of an angiogenesis polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino-or carboxyl-terminus of the angiogenesis polypeptide. The presence of such epitope-tagged forms of an angiogenesis polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the angiogenesis polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. In an alternative embodiment, the chimeric molecule may comprise a fusion of an angiogenesis polypeptide with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule, such a fusion could be to the Fc region of an IgG molecule.

Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; HIS6 and metal chelation tags, the flu HA tag polypeptide and its antibody 12CA5 [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the c-myc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al., Molecular and Cellular Biology, 5:3610-3616 (1985)];

and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

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Also included with an embodiment of angiogenesis protein are other angiogenesis proteins of the angiogenesis family, and angiogenesis proteins from other organisms, which are cloned and expressed as outlined below. Thus, probe or degenerate polymerase chain reaction (PCR) primer sequences may be used to find other related angiogenesis proteins from humans or other organisms. As will be appreciated by those in the art, particularly useful probe and/or PCR primer sequences include the unique areas of the angiogenesis nucleic acid sequence. As is generally known in the art, preferred PCR primers are from about 15 to about 35 nucleotides in length, with from about 20 to about 30 being preferred, and may contain inosine as needed. The conditions for the PCR reaction are well known in the art (e.g., Innis, PCR Protocols, supra).

In addition, as is outlined herein, angiogenesis proteins can be made that are longer than those encoded by the nucleic acids of the figures, e.g., by the elucidation of extended sequences, the addition of epitope or purification tags, the addition of other fusion sequences, etc.

Angiogenesis proteins may also be identified as being encoded by angiogenesis nucleic acids. Thus, angiogenesis proteins are encoded by nucleic acids that will hybridize to the sequences of the sequence listings, or their complements, as outlined herein.

In a preferred embodiment, when the angiogenesis protein is to be used to generate antibodies, e.g., for immunotherapy or immunodiagnosis, the angiogenesis protein should share at least one epitope or determinant with the full length protein. By "epitope" or "determinant" herein is typically meant a portion of a protein which will generate and/or bind an antibody or T-cell receptor in the context of MHC. Thus, in most instances, antibodies made to a smaller angiogenesis protein will be able to bind to the full-length protein, particularly linear epitopes. In a preferred embodiment, the epitope is unique; that is, antibodies generated to a unique epitope show little or no cross-reactivity. In a preferred embodiment, the epitope is selected from a protein sequence set out in Table 8.

Methods of preparing polyclonal antibodies are known to the skilled artisan (e.g., Coligan, supra; and Harlow & Lane, supra). Polyclonal antibodies can be raised in a mammal, e.g., by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include a protein encoded by a nucleic acid of the figures or fragment thereof or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

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The antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. The immunizing agent will typically include a polypeptide encoded by a nucleic acid of Tables 1-8, or fragment thereof, or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell [Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103]. Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

In one embodiment, the antibodies are bispecific antibodies. Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens or that have binding specificities for two epitopes on the same antigen. In one embodiment, one of the binding specificities is for a protein encoded by a nucleic acid Tables 1-8 or a fragment thereof, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit, preferably one that is tumor specific. Alternatively, tetramer-type technology may create multivalent reagents.

In a preferred embodiment, the antibodies to angiogenesis protein are capable of reducing or eliminating a biological function of an angiogenesis protein, as is described below. That is, the addition of anti-angiogenesis protein antibodies (either polyclonal or preferably monoclonal) to angiogenic tissue (or cells containing angiogenesis) may reduce or eliminate the angiogenesis activity. Generally, at least a 25% decrease in activity is preferred, with at least about 50% being particularly preferred and about a 95-100% decrease being especially preferred.

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In a preferred embodiment the antibodies to the angiogenesis proteins are humanized antibodies (e.g., Xenerex Biosciences, Mederex, Inc., Abgenix, Inc., Protein Design Labs, Inc.) Humanized forms of non-human (e.g., murine) antibodies are chimeric molecules of immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')2 or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues form a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a nonhuman species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, a humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the framework (FR) regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human

immunoglobulin [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-329 (1988); and Presta, Curr. Op. Struct. Biol., 2:593-596 (1992)].

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Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as import residues, which are typically taken from an import variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such humanized antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries [Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)]. The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985) and Boerner et al., J. Immunol., 147(1):86-95 (1991)]. Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10, 779-783 (1992); Lonberg et al., Nature 368 856-859 (1994); Morrison, Nature 368, 812-13 (1994); Fishwild et al., Nature Biotechnology 14, 845-51 (1996); Neuberger, Nature Biotechnology 14, 826 (1996); Lonberg and Huszar, Intern. Rev. Immunol. 13 65-93 (1995).

By immunotherapy is meant treatment of angiogenesis with an antibody raised against angiogenesis proteins. As used herein, immunotherapy can be passive or active. Passive immunotherapy as defined herein is the passive transfer of antibody to a recipient

(patient). Active immunization is the induction of antibody and/or T-cell responses in a recipient (patient). Induction of an immune response is the result of providing the recipient with an antigen to which antibodies are raised. As appreciated by one of ordinary skill in the art, the antigen may be provided by injecting a polypeptide against which antibodies are desired to be raised into a recipient, or contacting the recipient with a nucleic acid capable of expressing the antigen and under conditions for expression of the antigen, leading to an immune response.

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In a preferred embodiment the angiogenesis proteins against which antibodies are raised are secreted proteins as described above. Without being bound by theory, antibodies used for treatment, bind and prevent the secreted protein from binding to its receptor, thereby inactivating the secreted angiogenesis protein.

In another preferred embodiment, the angiogenesis protein to which antibodies are raised is a transmembrane protein. Without being bound by theory, antibodies used for treatment, bind the extracellular domain of the angiogenesis protein and prevent it from binding to other proteins, such as circulating ligands or cell-associated molecules. The antibody may cause down-regulation of the transmembrane angiogenesis protein. As will be appreciated by one of ordinary skill in the art, the antibody may be a competitive, noncompetitive or uncompetitive inhibitor of protein binding to the extracellular domain of the angiogenesis protein. The antibody is also an antagonist of the angiogenesis protein. Further, the antibody prevents activation of the transmembrane angiogenesis protein. In one aspect, when the antibody prevents the binding of other molecules to the angiogenesis protein, the antibody prevents growth of the cell. The antibody may also be used to target or sensitize the cell to cytotoxic agents, including, but not limited to TNF-α, TNF-β, IL-1, INF-γ and IL-2, or chemotherapeutic agents including 5FU, vinblastine, actinomycin D, cisplatin, methotrexate, and the like. In some instances the antibody belongs to a sub-type that activates serum complement when complexed with the transmembrane protein thereby mediating cytotoxicity or antigen-dependent cytotoxicity (ADCC). Thus, angiogenesis is treated by administering to a patient antibodies directed against the transmembrane angiogenesis protein. Antibody-labeling may activate a co-toxin, localize a toxin payload, or otherwise provide means to locally ablate cells.

In another preferred embodiment, the antibody is conjugated or fused to an effector moiety. The effector moiety can be any number of molecules, including labelling moieties such as radioactive labels or fluorescent labels, or can be a therapeutic moiety. In

one aspect the therapeutic moiety is a small molecule that modulates the activity of the angiogenesis protein. In another aspect the therapeutic moiety modulates the activity of molecules associated with or in close proximity to the angiogenesis protein. The therapeutic moiety may inhibit enzymatic activity such as protease or collagenase activity associated with angiogenesis, or be an attractant of other cells, such as NK cells.

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In a preferred embodiment, the therapeutic moiety can also be a cytotoxic agent. In this method, targeting the cytotoxic agent to angiogenesis tissue or cells, results in a reduction in the number of afflicted cells, thereby reducing symptoms associated with angiogenesis. Cytotoxic agents are numerous and varied and include, but are not limited to, cytotoxic drugs or toxins or active fragments of such toxins. Suitable toxins and their corresponding fragments include diphtheria A chain, exotoxin A chain, ricin A chain, abrin A chain, curcin, crotin, phenomycin, enomycin and the like. Cytotoxic agents also include radiochemicals made by conjugating radioisotopes to antibodies raised against angiogenesis proteins, or binding of a radionuclide to a chelating agent that has been covalently attached to the antibody. Targeting the therapeutic moiety to transmembrane angiogenesis proteins not only serves to increase the local concentration of therapeutic moiety in the angiogenesis afflicted area, but also serves to reduce deleterious side effects that may be associated with the therapeutic moiety.

In another preferred embodiment, the angiogenesis protein against which the antibodies are raised is an intracellular protein. In this case, the antibody may be conjugated or fused to a protein which facilitates entry into the cell. In one case, the antibody enters the cell by endocytosis. In another embodiment, a nucleic acid encoding the antibody is administered to the individual or cell. Moreover, wherein the angiogenesis protein can be targeted within a cell, i.e., the nucleus, an antibody thereto contains a signal for that target localization, i.e., a nuclear localization signal.

The angiogenesis antibodies of the invention specifically bind to angiogenesis proteins. By "specifically bind" herein is meant that the antibodies bind to the protein with a K_d of at least about 0.1 mM, more usually at least about 1 μ M, preferably at least about 0.1 μ M or better, and most preferably, 0.01 μ M or better. Selectivity of binding is also important.

In a preferred embodiment, the angiogenesis protein is purified or isolated after expression. Angiogenesis proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample. Standard purification methods include electrophoretic, molecular, immunological

and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the angiogenesis protein may be purified using a standard anti-angiogenesis protein antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes, R., Protein Purification, Springer-Verlag, NY (1982). The degree of purification necessary will vary depending on the use of the angiogenesis protein. In some instances no purification will be necessary.

Once expressed and purified if necessary, the angiogenesis proteins and nucleic acids are useful in a number of applications. They may be used as immunoselection reagents, as vaccine reagents, as screening agents, etc.

Detection of angiogenesis sequence for diagnostic and therapeutic applications

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In one aspect, the RNAexpression levels of genes are determined for different cellular states in the angiogenesis phenotype. Expression levels of genes in normal tissue (i.e., not undergoing angiogenesis) and in angiogenesis tissue (and in some cases, for varying severities of angiogenesis that relate to prognosis, as outlined below) are evaluated to provide expression profiles. An expression profile of a particular cell state or point of development is essentially a "fingerprint" of the state. While two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is reflective of the state of the cell. By comparing expression profiles of cells in different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. Then, diagnosis may be performed or confirmed to determine whether a tissue sample has the gene expression profile of normal or angiogenesic tissue. This will provide for molecular diagnosis of related conditions.

"Differential expression," or grammatical equivalents as used herein, refers to qualitative or quantitative differences in the temporal and/or cellular gene expression patterns within and among cells and tissue. Thus, a differentially expressed gene can qualitatively have its expression altered, including an activation or inactivation, in, e.g., normal versus angiogenic tissue. Genes may be turned on or turned off in a particular state, relative to another state thus permitting comparison of two or more statese. A qualitatively regulated gene will exhibit an expression pattern within a state or cell type which is detectable by standard techniques. Some genes will be expressed in one state or cell type, but not in both. Alternatively, the difference in expression may be quantitative, e.g., in that

expression is increased or decreased; *i.e.*, gene expression is either upregulated, resulting in an increased amount of transcript, or downregulated, resulting in a decreased amount of transcript. The degree to which expression differs need only be large enough to quantify via standard characterization techniques as outlined below, such as by use of Affymetrix

5 GeneChipTM expression arrays, Lockhart, Nature Biotechnology, 14:1675-1680 (1996), hereby expressly incorporated by reference. Other techniques include, but are not limited to, quantitative reverse transcriptase PCR, Northern analysis and RNase protection. As outlined above, preferably the change in expression (*i.e.*, upregulation or downregulation) is at least about 50%, more preferably at least about 100%, more preferably at least about 150%, more preferably at least about 200%, with from 300 to at least 1000% being especially preferred.

Evaluation may be at the gene transcript, or the protein level. The amount of gene expression may be monitored using nucleic acid probes to the DNA or RNA equivalent of the gene transcript, and the quantification of gene expression levels, or, alternatively, the final gene product itself (protein) can be monitored, e.g., with antibodies to the angiogenesis protein and standard immunoassays (ELISAs, etc.) or other techniques, including mass spectroscopy assays, 2D gel electrophoresis assays, etc. Proteins corresponding to angiogenesis genes, i.e., those identified as being important in an angiogenesis phenotype, can be evaluated in an angiogenesis diagnostic test.

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In a preferred embodiment, gene expression monitoring is performed simultaneously on a number of genes. Multiple protein expression monitoring can be performed as well. Similarly, these assays may be performed on an individual basis as well.

In this embodiment, the angiogenesis nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of angiogenesis sequences in a particular cell. The assays are further described below in the example. PCR techniques can be used to provide greater sensitivity.

In a preferred embodiment nucleic acids encoding the angiogenesis protein are detected. Although DNA or RNA encoding the angiogenesis protein may be detected, of particular interest are methods wherein an mRNA encoding an angiogenesis protein is detected. Probes to detect mRNA can be a nucleotide/deoxynucleotide probe that is complementary to and hybridizes with the mRNA and includes, but is not limited to, oligonucleotides, cDNA or RNA. Probes also should contain a detectable label, as defined herein. In one method the mRNA is detected after immobilizing the nucleic acid to be examined on a solid support such as nylon membranes and hybridizing the probe with the sample. Following washing to remove the non-specifically bound probe, the label is

detected. In another method detection of the mRNA is performed in situ. In this method permeabilized cells or tissue samples are contacted with a detectably labeled nucleic acid probe for sufficient time to allow the probe to hybridize with the target mRNA. Following washing to remove the non-specifically bound probe, the label is detected. For example a digoxygenin labeled riboprobe (RNA probe) that is complementary to the mRNA encoding an angiogenesis protein is detected by binding the digoxygenin with an anti-digoxygenin secondary antibody and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indoyl phosphate.

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In a preferred embodiment, various proteins from the three classes of proteins as described herein (secreted, transmembrane or intracellular proteins) are used in diagnostic assays. The angiogenesis proteins, antibodies, nucleic acids, modified proteins and cells containing angiogenesis sequences are used in diagnostic assays. This can be performed on an individual gene or corresponding polypeptide level. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes and/or corresponding polypeptides.

As described and defined herein, angiogenesis proteins, including intracellular, transmembrane or secreted proteins, find use as markers of angiogenesis. Detection of these proteins in putative angiogenesis tissue allows for detection or diagnosis of angiogenesis. In one embodiment, antibodies are used to detect angiogenesis proteins. A preferred method separates proteins from a sample by electrophoresis on a gel (typically a denaturing and reducing protein gel, but may be another type of gel, including isoelectric focusing gels and the like). Following separation of proteins, the angiogenesis protein is detected, e.g., by immunoblotting with antibodies raised against the angiogenesis protein. Methods of immunoblotting are well known to those of ordinary skill in the art.

In another preferred method, antibodies to the angiogenesis protein find use in in situ imaging techniques, e.g., in histology (e.g., Methods in Cell Biology: Antibodies in Cell Biology, volume 37 (Asai, ed. 1993)). In this method cells are contacted with from one to many antibodies to the angiogenesis protein(s). Following washing to remove non-specific antibody binding, the presence of the antibody or antibodies is detected. In one embodiment the antibody is detected by incubating with a secondary antibody that contains a detectable label. In another method the primary antibody to the angiogenesis protein(s) contains a detectable label, for example an enzyme marker that can act on a substrate. In another preferred embodiment each one of multiple primary antibodies contains a distinct and

detectable label. This method finds particular use in simultaneous screening for a plurality of angiogenesis proteins. As will be appreciated by one of ordinary skill in the art, many other histological imaging techniques are also provided by the invention.

In a preferred embodiment the label is detected in a fluorometer which has the ability to detect and distinguish emissions of different wavelengths. In addition, a fluorescence activated cell sorter (FACS) can be used in the method.

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In another preferred embodiment, antibodies find use in diagnosing angiogenesis from biological samples, such as blood, urine, sputum, or other bodily fluids. As previously described, certain angiogenesis proteins are secreted/circulating molecules. Blood samples, therefore, are useful as samples to be probed or tested for the presence of secreted angiogenesis proteins. Antibodies can be used to detect an angiogenesis protein by previously described immunoassay techniques including ELISA, immunoblotting (Western blotting), immunoprecipitation, BIACORE technology and the like. Conversely, the presence of antibodies may indicate an immune response against an endogenous angiogenesis protein.

In a preferred embodiment, in situ hybridization of labeled angiogenesis nucleic acid probes to tissue arrays is done. For example, arrays of tissue samples, including angiogenesis tissue and/or normal tissue, are made. In situ hybridization (see, e.g., Ausubel, supra) is then performed. When comparing the fingerprints between an individual and a standard, the skilled artisan can make a diagnosis, a prognosis, or a prediction based on the findings. It is further understood that the genes which indicate the diagnosis may differ from those which indicate the prognosis and molecular profiling of the condition of the cells may lead to distinctions between responsive or refractory conditions or may be predictive of outcomes.

In a preferred embodiment, the angiogenesis proteins, antibodies, nucleic acids, modified proteins and cells containing angiogenesis sequences are used in prognosis assays. As above, gene expression profiles can be generated that correlate to angiogenesis severity, in terms of long term prognosis. Again, this may be done on either a protein or gene level, with the use of genes being preferred. As above, angiogenesis probes may be attached to biochips for the detection and quantification of angiogenesis sequences in a tissue or patient. The assays proceed as outlined above for diagnosis. PCR method may provide more sensitive and accurate quantification.

In a preferred embodiment members of the three classes of proteins as described herein are used in drug screening assays. The angiogenesis proteins, antibodies, nucleic acids, modified proteins and cells containing angiogenesis sequences are used in drug

screening assays or by evaluating the effect of drug candidates on a "gene expression profile" or expression profile of polypeptides. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent (e.g., Zlokarnik, et al., Science 279, 84-8 (1998); Heid, Genome Res 6:986-94, 1996).

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In a preferred embodiment, the angiogenesis proteins, antibodies, nucleic acids, modified proteins and cells containing the native or modified angiogenesis proteins are used in screening assays. That is, the present invention provides novel methods for screening for compositions which modulate the angiogenesis phenotype or an identified physiological function of an angiogenesis protein. As above, this can be done on an individual gene level or by evaluating the effect of drug candidates on a "gene expression profile". In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, see Zlokarnik, *supra*.

Having identified the differentially expressed genes herein, a variety of assays may be executed. In a preferred embodiment, assays may be run on an individual gene or protein level. That is, having identified a particular gene as up regulated in angiogenesis, test compounds can be screened for the ability to modulate gene expression or for binding to the angiogenic protein. "Modulation" thus includes both an increase and a decrease in gene expression. The preferred amount of modulation will depend on the original change of the gene expression in normal versus tissue undergoing angiogenesis, with changes of at least 10%, preferably 50%, more preferably 100-300%, and in some embodiments 300-1000% or greater. Thus, if a gene exhibits a 4-fold increase in angiogenic tissue compared to normal tissue, a decrease of about four-fold is often desired; similarly, a 10-fold decrease in angiogenic tissue compared to normal tissue often provides a target value of a 10-fold increase in expression to be induced by the test compound.

The amount of gene expression may be monitored using nucleic acid probes and the quantification of gene expression levels, or, alternatively, the gene product itself can be monitored, e.g., through the use of antibodies to the angiogenesis protein and standard immunoassays. Proteomics and separation techniques may also allow quantification of expression.

In a preferred embodiment, gene expression or protein monitoring of a number of entitites, *i.e.*, an expression profile, is monitored simultaneously. Such profiles will typically invove a plurality of those entitites described herein..

In this embodiment, the angiogenesis nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of angiogenesis sequences in a particular cell. Alternatively, PCR may be used. Thus, a series, e.g., of microtiter plate, may be used with dispensed primers in desired wells. A PCR reaction can then be performed and analyzed for each well.

Modulators of angiogenesis

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Expression monitoring can be performed to identify compounds that modify the expression of one or more angiogenesis-associated sequences, e.g., a polynucleotide sequence set out in Tables 1-8. Generally, in a preferred embodiment, a test modulator is added to the cells prior to analysis. Moreover, screens are also provided to identify agents that modulate angiogenesis, modulate angiogenesis proteins, bind to an angiogenesis protein, or interfere with the binding of an angiogenesis protein and an antibody or other binding partner.

The term "test compound" or "drug candidate" or "modulator" or grammatical equivalents as used herein describes any molecule, e.g., protein, oligopeptide, small organic molecule, polysaccharide, polynucleotide, etc., to be tested for the capacity to directly or indirectly alter the angiogenesis phenotype or the expression of an angiogenesis sequence, e.g., a nucleic acid or protein sequence. In preferred embodiments, modulators alter expression profiles, or expression profile nucleic acids or proteins provided herein. In one embodiment, the modulator suppresses an angiogenesis phenotype, for example to a normal tissue fingerprint. In another embodiment, a modulator induced an angiogenesis phenotype. Generally, a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, i.e., at zero concentration or below the level of detection.

In one aspect, a modulator will neutralize the effect of an angiogenesis protein. By "neutralize" is meant that activity of a protein is inhibited or blocked and thereby has substantially no effect on a cell.

In certain embodiments, combinatorial libraries of potential modulators will be screened for an ability to bind to an angiogenesis polypeptide or to modulate activity. Conventionally, new chemical entities with useful properties are generated by identifying a chemical compound (called a "lead compound") with some desirable property or activity, e.g., inhibiting activity, creating variants of the lead compound, and evaluating the property

and activity of those variant compounds. Often, high throughput screening (HTS) methods are employed for such an analysis.

In one preferred embodiment, high throughput screening methods involve providing a library containing a large number of potential therapeutic compounds (candidate compounds). Such "combinatorial chemical libraries" are then screened in one or more assays to identify those library members (particular chemical species or subclasses) that display a desired characteristic activity. The compounds thus identified can serve as conventional "lead compounds" or can themselves be used as potential or actual therapeutics.

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A combinatorial chemical library is a collection of diverse chemical compounds generated by either chemical synthesis or biological synthesis by combining a number of chemical "building blocks" such as reagents. For example, a linear combinatorial chemical library, such as a polypeptide (e.g., mutein) library, is formed by combining a set of chemical building blocks called amino acids in every possible way for a given compound length (i.e., the number of amino acids in a polypeptide compound). Millions of chemical compounds can be synthesized through such combinatorial mixing of chemical building blocks (Gallop et al. (1994) J. Med. Chem. 37(9): 1233-1251).

Preparation and screening of combinatorial chemical libraries is well known to those of skill in the art. Such combinatorial chemical libraries include, but are not limited to, peptide libraries (see, e.g., U.S. Patent No. 5,010,175, Furka (1991) Int. J. Pept. Prot. Res., 37: 487-493, Houghton et al. (1991) Nature, 354: 84-88), peptoids (PCT Publication No WO 91/19735, 26 Dec. 1991), encoded peptides (PCT Publication WO 93/20242, 14 Oct. 1993), random bio-oligomers (PCT Publication WO 92/00091, 9 Jan. 1992), benzodiazepines (U.S. Pat. No. 5,288,514), diversomers such as hydantoins, benzodiazepines and dipeptides (Hobbs et al., (1993) Proc. Nat. Acad. Sci. USA 90: 6909-6913), vinylogous polypeptides (Hagihara et al. (1992) J. Amer. Chem. Soc. 114: 6568), nonpeptidal peptidomimetics with a Beta-D-Glucose scaffolding (Hirschmann et al., (1992) J. Amer. Chem. Soc. 114: 9217-9218), analogous organic syntheses of small compound libraries (Chen et al. (1994) J. Amer. Chem. Soc. 116: 2661), oligocarbamates (Cho, et al., (1993) Science 261:1303), and/or peptidyl phosphonates (Campbell et al., (1994) J. Org. Chem. 59: 658). See, generally, Gordon et al., (1994) J. Med. Chem. 37:1385, nucleic acid libraries (see, e.g., Strategene, Corp.), peptide nucleic acid libraries (see, e.g., U.S. Patent 5,539,083), antibody libraries (see, e.g., Vaughn et al. (1996) Nature Biotechnology, 14(3): 309-314), and PCT/US96/10287), carbohydrate libraries (see, e.g., Liang et al., (1996) Science, 274: 1520-1522, and U.S. Patent No. 5,593,853), and small organic molecule libraries (see, e.g., benzodiazepines, Baum (1993)

C&EN, Jan 18, page 33; isoprenoids, U.S. Patent No. 5,569,588; thiazolidinones and metathiazanones, U.S. Patent No. 5,549,974; pyrrolidines, U.S. Patent Nos. 5,525,735 and 5,519,134; morpholino compounds, U.S. Patent No. 5,506,337; benzodiazepines, U.S. Patent No. 5,288,514; and the like).

Devices for the preparation of combinatorial libraries are commercially available (see, e.g., 357 MPS, 390 MPS, Advanced Chem Tech, Louisville KY, Symphony, Rainin, Woburn, MA, 433A Applied Biosystems, Foster City, CA, 9050 Plus, Millipore, Bedford, MA).

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A number of well known robotic systems have also been developed for solution phase chemistries. These systems include automated workstations like the automated synthesis apparatus developed by Takeda Chemical Industries, LTD. (Osaka, Japan) and many robotic systems utilizing robotic arms (Zymate II, Zymark Corporation, Hopkinton, Mass.; Orca, Hewlett-Packard, Palo Alto, Calif.), which mimic the manual synthetic operations performed by a chemist. Any of the above devices are suitable for use with the present invention. The nature and implementation of modifications to these devices (if any) so that they can operate as discussed herein will be apparent to persons skilled in the relevant art. In addition, numerous combinatorial libraries are themselves commercially available (see, e.g., ComGenex, Princeton, N.J., Asinex, Moscow, Ru, Tripos, Inc., St. Louis, MO, ChemStar, Ltd, Moscow, RU, 3D Pharmaceuticals, Exton, PA, Martek Biosciences, Columbia, MD, etc.).

The assays to identify modulators are amenable to high throughput screening. Preferred assays thus detect enhancement or inhibition of angiogenesis gene transcription, inhibition or enhancement of polypeptide expression, and inhibition or enhancement of polypeptide activity.

High throughput assays for the presence, absence, quantification, or other properties of particular nucleic acids or protein products are well known to those of skill in the art. Similarly, binding assays and reporter gene assays are similarly well known. Thus, for example, U.S. Patent No. 5,559,410 discloses high throughput screening methods for proteins, U.S. Patent No. 5,585,639 discloses high throughput screening methods for nucleic acid binding (i.e., in arrays), while U.S. Patent Nos. 5,576,220 and 5,541,061 disclose high throughput methods of screening for ligand/antibody binding.

In addition, high throughput screening systems are commercially available (see, e.g., Zymark Corp., Hopkinton, MA; Air Technical Industries, Mentor, OH; Beckman Instruments, Inc. Fullerton, CA; Precision Systems, Inc., Natick, MA, etc.). These systems

typically automate entire procedures, including all sample and reagent pipetting, liquid dispensing, timed incubations, and final readings of the microplate in detector(s) appropriate for the assay. These configurable systems provide high throughput and rapid start up as well as a high degree of flexibility and customization. The manufacturers of such systems provide detailed protocols for various high throughput systems. Thus, for example, Zymark Corp. provides technical bulletins describing screening systems for detecting the modulation of gene transcription, ligand binding, and the like.

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In one embodiment, modulators are proteins, often naturally occurring proteins or fragments of naturally occurring proteins. Thus, e.g., cellular extracts containing proteins, or random or directed digests of proteinaceous cellular extracts, may be used. In this way libraries of proteins may be made for screening in the methods of the invention. Particularly preferred in this embodiment are libraries of bacterial, fungal, viral, and mammalian proteins, with the latter being preferred, and human proteins being especially preferred. Paticularly useful test compound will be directed to the class of proteins to which the target belongs, e.g., substrates for enzymes or ligands and receptors.

In a preferred embodiment, modulators are peptides of from about 5 to about 30 amino acids, with from about 5 to about 20 amino acids being preferred, and from about 7 to about 15 being particularly preferred. The peptides may be digests of naturally occurring proteins as is outlined above, random peptides, or "biased" random peptides. By "randomized" or grammatical equivalents herein is meant that each nucleic acid and peptide consists of essentially random nucleotides and amino acids, respectively. Since generally these random peptides (or nucleic acids, discussed below) are chemically synthesized, they may incorporate any nucleotide or amino acid at any position. The synthetic process can be designed to generate randomized proteins or nucleic acids, to allow the formation of all or most of the possible combinations over the length of the sequence, thus forming a library of randomized candidate bioactive proteinaceous agents.

In one embodiment, the library is fully randomized, with no sequence preferences or constants at any position. In a preferred embodiment, the library is biased. That is, some positions within the sequence are either held constant, or are selected from a limited number of possibilities. For example, in a preferred embodiment, the nucleotides or amino acid residues are randomized within a defined class, for example, of hydrophobic amino acids, hydrophilic residues, sterically biased (either small or large) residues, towards the creation of nucleic acid binding domains, the creation of cysteines, for cross-linking,

prolines for SH-3 domains, serines, threonines, tyrosines or histidines for phosphorylation sites, etc., or to purines, etc.

Modulators of angiogenesis can also be nucleic acids, as defined above.

As described above generally for proteins, nucleic acid modulating agents may be naturally occurring nucleic acids, random nucleic acids, or "biased" random nucleic acids. For example, digests of procaryotic or eucaryotic genomes may be used as is outlined above for proteins.

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In a preferred embodiment, the candidate compounds are organic chemical moieties, a wide variety of which are available in the literature.

After the candidate agent has been added and the cells allowed to incubate for some period of time, the sample containing a target sequence to be analyzed is added to the biochip. If required, the target sequence is prepared using known techniques. For example, the sample may be treated to lyse the cells, using known lysis buffers, electroporation, etc., with purification and/or amplification such as PCR performed as appropriate. For example, an *in vitro* transcription with labels covalently attached to the nucleotides is performed. Generally, the nucleic acids are labeled with biotin-FITC or PE, or with cy3 or cy5.

In a preferred embodiment, the target sequence is labeled with, for example, a fluorescent, a chemiluminescent, a chemical, or a radioactive signal, to provide a means of detecting the target sequence's specific binding to a probe. The label also can be an enzyme, such as, alkaline phosphatase or horseradish peroxidase, which when provided with an appropriate substrate produces a product that can be detected. Alternatively, the label can be a labeled compound or small molecule, such as an enzyme inhibitor, that binds but is not catalyzed or altered by the enzyme. The label also can be a moiety or compound, such as, an epitope tag or biotin which specifically binds to streptavidin. For the example of biotin, the streptavidin is labeled as described above, thereby, providing a detectable signal for the bound target sequence. Unbound labeled streptavidin is typically removed prior to analysis.

As will be appreciated by those in the art, these assays can be direct hybridization assays or can comprise "sandwich assays", which include the use of multiple probes, as is generally outlined in U.S. Patent Nos. 5,681,702, 5,597,909, 5,545,730, 5,594,117, 5,591,584, 5,571,670, 5,580,731, 5,571,670, 5,591,584, 5,624,802, 5,635,352, 5,594,118, 5,359,100, 5,124,246 and 5,681,697, all of which are hereby incorporated by reference. In this embodiment, in general, the target nucleic acid is prepared as outlined above, and then added to the biochip comprising a plurality of nucleic acid probes, under conditions that allow the formation of a hybridization complex.

A variety of hybridization conditions may be used in the present invention, including high, moderate and low stringency conditions as outlined above. The assays are generally run under stringency conditions which allows formation of the label probe hybridization complex only in the presence of target. Stringency can be controlled by altering a step parameter that is a thermodynamic variable, including, but not limited to, temperature, formamide concentration, salt concentration, chaotropic salt concentration pH, organic solvent concentration, etc.

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These parameters may also be used to control non-specific binding, as is generally outlined in U.S. Patent No. 5,681,697. Thus it may be desirable to perform certain steps at higher stringency conditions to reduce non-specific binding.

The reactions outlined herein may be accomplished in a variety of ways. Components of the reaction may be added simultaneously, or sequentially, in different orders, with preferred embodiments outlined below. In addition, the reaction may include a variety of other reagents. These include salts, buffers, neutral proteins, e.g. albumin, detergents, etc. which may be used to facilitate optimal hybridization and detection, and/or reduce non-specific or background interactions. Reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may also be used as appropriate, depending on the sample preparation methods and purity of the target.

The assay data are analyzed to determine the expression levels, and changes in expression levels as between states, of individual genes, forming a gene expression profile.

Screens are performed to identify modulators of the angiogenesis phenotype. In one embodiment, screening is performed to identify modulators that can induce or suppress a particular expression profile, thus preferably generating the associated phenotype. In another embodiment, e.g., for diagnostic applications, having identified differentially expressed genes important in a particular state, screens can be performed to identify modulators that alter expression of individual genes. In an another embodiment, screening is performed to identify modulators that alter a biological function of the expression product of a differentially expressed gene. Again, having identified the importance of a gene in a particular state, screens are performed to identify agents that bind and/or modulate the biological activity of the gene product.

In addition screens can be done for genes that are induced in response to a candidate agent. After identifying a modulator based upon its ability to suppress an angiogenesis expression pattern leading to a normal expression pattern, or to modulate a single angiogenesis gene expression profile so as to mimic the expression of the gene from

normal tissue, a screen as described above can be performed to identify genes that are specifically modulated in response to the agent. Comparing expression profiles between normal tissue and agent treated angiogenesis tissue reveals genes that are not expressed in normal tissue or angiogenesis tissue, but are expressed in agent treated tissue. These agent-specific sequences can be identified and used by methods described herein for angiogenesis genes or proteins. In particular these sequences and the proteins they encode find use in marking or identifying agent treated cells. In addition, antibodies can be raised against the agent induced proteins and used to target novel therapeutics to the treated angiogenesis tissue sample.

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Thus, in one embodiment, a test compound is administered to a population of angiogenic cells, that have an associated angiogenesis expression profile. By "administration" or "contacting" herein is meant that the candidate agent is added to the cells in such a manner as to allow the agent to act upon the cell, whether by uptake and intracellular action, or by action at the cell surface. In some embodiments, nucleic acid encoding a proteinaceous candidate agent (i.e., a peptide) may be put into a viral construct such as an adenoviral or retroviral construct, and added to the cell, such that expression of the peptide agent is accomplished, e.g., PCT US97/01019. Regulatable gene therapy systems can also be used.

Once the test compound has been administered to the cells, the cells can be washed if desired and are allowed to incubate under preferably physiological conditions for some period of time. The cells are then harvested and a new gene expression profile is generated, as outlined herein.

Thus, for example, angiogenesis tissue may be screened for agents that modulate, e.g., induce or suppress the angiogenesis phenotype. A change in at least one gene, preferably many, of the expression profile indicates that the agent has an effect on angiogenesis activity. By defining such a signature for the angiogenesis phenotype, screens for new drugs that alter the phenotype can be devised. With this approach, the drug target need not be known and need not be represented in the original expression screening platform, nor does the level of transcript for the target protein need to change.

Measure of angiogenesis polypeptide activity, or of angiogenesis or the angiogenic phenotype can be performed using a variety of assays. For example, the effects of the test compounds upon the function of the anagiogenesis polypeptides can be measured by examining parameters described above. A suitable physiological change that affects activity can be used to assess the influence of a test compound on the polypeptides of this invention.

When the functional consequences are determined using intact cells or animals, one can also measure a variety of effects such as, in the case of angiogenesis associated with tumors, tumor growth, neovascularization, hormone release, transcriptional changes to both known and uncharacterized genetic markers (e.g., northern blots), changes in cell metabolism such as cell growth or pH changes, and changes in intracellular second messengers such as cGMP. In the assays of the invention, mammalian angiogenesis polypeptide is typically used, e.g., mouse, preferably human.

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A variety of angiogenesis assays are known to those of skill in the art. Various models have been employed to evaluate angiogenesis (e.g., Croix et al., Science 289:1197-1202, 2000 and Kahn et al., Amer. J. Pathol. 156:1887-1900). Assessement of angiogenesis in the presence of a potential modulator of angiogenesis can be performed using cell-cultrebased angiogenesis assays, e.g., endothelial cell tube formation assays, as well as other bioassays such as the chick CAM assay, the mouse corneal assay, and assays measuring the effect of administering potential modulators on implanted tumors. The chick CAM assay is described by O'Reilly, et al. Cell 79: 315-328, 1994. Briefly, 3 day old chicken embryos with intact yolks are separated from the egg and placed in a petri dish. After 3 days of incubation, a methylcellulose disc containing the protein to be tested is applied to the CAM of individual embryos. After about 48 hours of incubation, the embryos and CAMs are observed to determine whether endothelial growth has been inhibited. The mouse corneal assay involves implanting a growth factor-containing pellet, along with another pellet containing the suspected endothelial growth inhibitor, in the comea of a mouse and observing the pattern of capillaries that are elaborated in the comea. Angiogenesis can also be measured by determining the extent of neovascularization of a tumor. For example, carcinoma cells can be subcutaneously inoculated into athymic nude mice and tumor growth then monitored. The cancer cells are treated with an angiogenesis inhibitor, such as an antibody, or other compound that is exogenously administered, or can be transfected prior to inoculation with a polynucleotide inhibitor of angiogenesis. Immunoassays using endothelial cell-specific antibodies are typically used to stain for vascularization of tumor and the number of vessels in the tumor.

Assays to identify compounds with modulating activity can be performed in vitro. For example, an angiogenesis polypeptide is first contacted with a potential modulator and incubated for a suitable amount of time, e.g., from 0.5 to 48 hours. In one embodiment, the angiogenesis polypeptide levels are determined in vitro by measuring the level of protein or mRNA. The level of protein is measured using immunoassays such as western blotting,

ELISA and the like with an antibody that selectively binds to the angiogenesis polypeptide or a fragment thereof. For measurement of mRNA, amplification, e.g., using PCR, LCR, or hybridization assays, e.g., northern hybridization, RNAse protection, dot blotting, are preferred. The level of protein or mRNA is detected using directly or indirectly labeled detection agents, e.g., fluorescently or radioactively labeled nucleic acids, radioactively or enzymatically labeled antibodies, and the like, as described herein.

Alternatively, a reporter gene system can be devised using the angiogenesis protein promoter operably linked to a reporter gene such as luciferase, green fluorescent protein, CAT, or β -gal. The reporter construct is typically transfected into a cell. After treatment with a potential modulator, the amount of reporter gene transcription, translation, or activity is measured according to standard techniques known to those of skill in the art.

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In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "angiogenesis proteins". In preferred embodiments the angiogenesis protein comprises a sequence shown in Table 8. The angiogenesis protein may be a fragment, or alternatively, be the full length protein to a fragment shown herein.

Preferably, the angiogenesis protein is a fragment of approximately 14 to 24 amino acids long. More preferably the fragment is a soluble fragment. In one embodiment an angiogenesis protein is conjugated or fused to an immunogenic agent or BSA.

In one embodiment, screening for modulators of expression of specific genes is performed. Typically, the expression of only one or a few genes are evaluated. In another embodiment, screens are designed to first find compounds that bind to differentially expressed proteins. These compounds are then evaluated for the ability to modulate differentially expressed activity. Moreover, once initial candidate compounds are identified, variants can be further screened to better evaluate structure activity relationships.

In a preferred embodiment, binding assays are done. In general, purified or isolated gene product is used; that is, the gene products of one or more differentially expressed nucleic acids are made. For example, antibodies are generated to the protein gene products, and standard immunoassays are run to determine the amount of protein present. Alternatively, cells comprising the angiogenesis proteins can be used in the assays.

Thus, in a preferred embodiment, the methods comprise combining an angiogenesis protein and a candidate compound, and determining the binding of the compound to the angiogenesis protein. Preferred embodiments utilize the human angiogenesis protein, although other mammalian proteins may also be used, for example for the development of animal models of human disease. In some embodiments, as outlined herein, variant or derivative angiogenesis proteins may be used.

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Generally, in a preferred embodiment of the methods herein, the angiogenesis protein or the candidate agent is non-diffusably bound to an insoluble support having isolated sample receiving areas (e.g. a microtiter plate, an array, etc.). The insoluble supports may be made of any composition to which the compositions can be bound, is readily separated from soluble material, and is otherwise compatible with the overall method of screening. The surface of such supports may be solid or porous and of any convenient shape. Examples of suitable insoluble supports include microtiter plates, arrays, membranes and beads. These are typically made of glass, plastic (e.g., polystyrene), polysaccharides, nylon or nitrocellulose, teflon™, etc. Microtiter plates and arrays are especially convenient because a large number of assays can be carried out simultaneously, using small amounts of reagents and samples. The particular manner of binding of the composition is not crucial so long as it is compatible with the reagents and overall methods of the invention, maintains the activity of the composition and is nondiffusable. Preferred methods of binding include the use of antibodies (which do not sterically block either the ligand binding site or activation sequence when the protein is bound to the support), direct binding to "sticky" or ionic supports, chemical crosslinking, the synthesis of the protein or agent on the surface, etc. Following binding of the protein or agent, excess unbound material is removed by washing. The sample receiving areas may then be blocked through incubation with bovine serum albumin (BSA), casein or other innocuous protein or other moiety.

In a preferred embodiment, the angiogenesis protein is bound to the support, and a test compound is added to the assay. Alternatively, the candidate agent is bound to the support and the angiogenesis protein is added. Novel binding agents include specific antibodies, non-natural binding agents identified in screens of chemical libraries, peptide analogs, etc. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including labeled in vitro protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, functional assays (phosphorylation assays, etc.) and the like.

The determination of the binding of the test modulating compound to the angiogenesis protein may be done in a number of ways. In a preferred embodiment, the compound is labelled, and binding determined directly, e.g., by attaching all or a portion of the angiogenesis protein to a solid support, adding a labelled candidate agent (e.g., a fluorescent label), washing off excess reagent, and determining whether the label is present on the solid support. Various blocking and washing steps may be utilized as appropriate.

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By "labeled" herein is meant that the compound is either directly or indirectly labeled with a label which provides a detectable signal, e.g. radioisotope, fluorescers, enzyme, antibodies, particles such as magnetic particles, chemiluminescers, or specific binding molecules, etc. Specific binding molecules include pairs, such as biotin and streptavidin, digoxin and antidigoxin, etc. For the specific binding members, the complementary member would normally be labeled with a molecule which provides for detection, in accordance with known procedures, as outlined above. The label can directly or indirectly provide a detectable signal.

In some embodiments, only one of the components is labeled, e.g., the proteins (or proteinaceous candidate compounds) can be labeled. Alternatively, more than one component can be labeled with different labels, e.g., ¹²⁵I for the proteins and a fluorophor for the compound. Proximity reagents, e.g., quenching or energy transfer reagents are also useful.

In one embodiment, the binding of the test compound is determined by competitive binding assay. The competitor is a binding moiety known to bind to the target molecule (*i.e.* an angiogenesis protein), such as an antibody, peptide, binding partner, ligand, etc. Under certain circumstances, there may be competitive binding between the compound and the binding moiety, with the binding moiety displacing the compound. In one embodiment, the test compound is labeled. Either the compound, or the competitor, or both, is added first to the protein for a time sufficient to allow binding, if present. Incubations may be performed at a temperature which facilitates optimal activity, typically between 4 and 40°C. Incubation periods are typically optimized, *e.g.*, to facilitate rapid high throughput screening. Typically between 0.1 and 1 hour will be sufficient. Excess reagent is generally removed or washed away. The second component is then added, and the presence or absence of the labeled component is followed, to indicate binding.

In a preferred embodiment, the competitor is added first, followed by the test compound. Displacement of the competitor is an indication that the test compound is binding to the angiogenesis protein and thus is capable of binding to, and potentially modulating, the

activity of the angiogenesis protein. In this embodiment, either component can be labeled. Thus, for example, if the competitor is labeled, the presence of label in the wash solution indicates displacement by the agent. Alternatively, if the test compound is labeled, the presence of the label on the support indicates displacement.

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In an alternative embodiment, the test compound is added first, with incubation and washing, followed by the competitor. The absence of binding by the competitor may indicate that the test compound is bound to the angiogenesis protein with a higher affinity. Thus, if the test compound is labeled, the presence of the label on the support, coupled with a lack of competitor binding, may indicate that the test compound is capable of binding to the angiogenesis protein.

In a preferred embodiment, the methods comprise differential screening to identity agents that are capable of modulating the activitity of the angiogenesis proteins. In this embodiment, the methods comprise combining an angiogenesis protein and a competitor in a first sample. A second sample comprises a test compound, an angiogenesis protein, and a competitor. The binding of the competitor is determined for both samples, and a change, or difference in binding between the two samples indicates the presence of an agent capable of binding to the angiogenesis protein and potentially modulating its activity. That is, if the binding of the competitor is different in the second sample relative to the first sample, the agent is capable of binding to the angiogenesis protein.

Alternatively, differential screening is used to identify drug candidates that bind to the native angiogenesis protein, but cannot bind to modified angiogenesis proteins. The structure of the angiogenesis protein may be modeled, and used in rational drug design to synthesize agents that interact with that site. Drug candidates that affect the activity of an angiogenesis protein are also identified by screening drugs for the ability to either enhance or reduce the activity of the protein.

Positive controls and negative controls may be used in the assays. Preferably control and test samples are performed in at least triplicate to obtain statistically significant results. Incubation of all samples is for a time sufficient for the binding of the agent to the protein. Following incubation, samples are washed free of non-specifically bound material and the amount of bound, generally labeled agent determined. For example, where a radiolabel is employed, the samples may be counted in a scintillation counter to determine the amount of bound compound.

A variety of other reagents may be included in the screening assays. These include reagents like salts, neutral proteins, e.g. albumin, detergents, etc. which may be used

to facilitate optimal protein-protein binding and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used. The mixture of components may be added in an order that provides for the requisite binding.

In a preferred embodiment, the invention provides methods for screening for a compound capable of modulating the activity of an angiogenesis protein. The methods comprise adding a test compound, as defined above, to a cell comprising angiogenesis proteins. Preferred cell types include almost any cell. The cells contain a recombinant nucleic acid that encodes an angiogenesis protein. In a preferred embodiment, a library of

candidate agents are tested on a plurality of cells.

In one aspect, the assays are evaluated in the presence or absence or previous or subsequent exposure of physiological signals, for example hormones, antibodies, peptides, antigens, cytokines, growth factors, action potentials, pharmacological agents including chemotherapeutics, radiation, carcinogenics, or other cells (i.e. cell-cell contacts). In another example, the determinations are determined at different stages of the cell cycle process.

In this way, compounds that modulate angiogenesis agents are identified. Compounds with pharmacological activity are able to enhance or interfere with the activity of the angiogenesis protein. Once identified, similar structures are evaluated to identify critical structural feature of the compound.

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In one embodiment, a method of inhibiting angiogenic cell division is provided. The method comprises administration of an angiogenesis inhibitor. In another embodiment, a method of inhibiting angiogenesis is provided. The method comprises administration of an angiogenesis inhibitor. In a further embodiment, methods of treating cells or individuals with angiogenesis are provided. The method comprises administration of an angiogenesis inhibitor.

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In one embodiment, an angiogenesis inhibitor is an antibody as discussed above. In another embodiment, the angiogenesis inhibitor is an antisense molecule.

Polynucleotide modulators of angiogenesis

Antisense Polynucleotides

In certain embodiments, the activity of an angiogenesis-associated protein is downregulated, or entirely inhibited, by the use of antisense polynucleotide, *i.e.*, a nucleic acid complementary to, and which can preferably hybridize specifically to, a coding mRNA nucleic acid sequence, *e.g.*, an angiogenesis protein mRNA, or a subsequence thereof.

Binding of the antisense polynucleotide to the mRNA reduces the translation and/or stability of the mRNA.

In the context of this invention, antisense polynucleotides can comprise naturally-occurring nucleotides, or synthetic species formed from naturally-occurring subunits or their close homologs. Antisense polynucleotides may also have altered sugar moieties or inter-sugar linkages. Exemplary among these are the phosphorothioate and other sulfur containing species which are known for use in the art. Analogs are comprehended by this invention so long as they function effectively to hybridize with the angiogenesis protein mRNA. See, e.g., Isis Pharmaceuticals, Carlsbad, CA; Sequitor, Inc., Natick, MA.

Such antisense polynucleotides can readily be synthesized using recombinant means, or can be synthesized *in vitro*. Equipment for such synthesis is sold by several vendors, including Applied Biosystems. The preparation of other oligonucleotides such as phosphorothioates and alkylated derivatives is also well known to those of skill in the art.

Antisense molecules as used herein include antisense or sense oligonucleotides. Sense oligonucleotides can, e.g., be employed to block trancription by binding to the anti-sense strand. The antisense and sense oligonucleotide comprise a single-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target mRNA (sense) or DNA (antisense) sequences for angiogenesis molecules. A preferred antisense molecule is for an angiogenesis sequences in Tables 1-8, or for a ligand or activator thereof. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment generally at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, for example, Stein and Cohen (Cancer Res. 48:2659, 1988) and van der Krol et al. (BioTechniques 6:958, 1988).

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Ribozymes

In addition to antisense polynucleotides, ribozymes can be used to target and inhibit transcription of angiogenesis-associated nucleotide sequences. A ribozyme is an RNA molecule that catalytically cleaves other RNA molecules. Different kinds of ribozymes have been described, including group I ribozymes, hammerhead ribozymes, hairpin ribozymes, RNase P, and axhead ribozymes (see, e.g., Castanotto et al. (1994) Adv. in Pharmacology 25: 289-317 for a general review of the properties of different ribozymes).

The general features of hairpin ribozymes are described, e.g., in Hampel et al. (1990) Nucl. Acids Res. 18: 299-304; Hampel et al. (1990) European Patent Publication No. 0

360 257; U.S. Patent No. 5,254,678. Methods of preparing are well known to those of skill in the art (see, e.g., Wong-Staal et al., WO 94/26877; Ojwang et al. (1993) Proc. Natl. Acad. Sci. USA 90: 6340-6344; Yamada et al. (1994) Human Gene Therapy 1: 39-45; Leavitt et al. (1995) Proc. Natl. Acad. Sci. USA 92: 699-703; Leavitt et al. (1994) Human Gene Therapy 5: 1151-120; and Yamada et al. (1994) Virology 205: 121-126).

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Polynucleotide modulators of angiogenesis may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell. Alternatively, a polynucleotide modulator of angiogenesis may be introduced into a cell containing the target nucleic acid sequence, e.g., by formation of an polynucleotide-lipid complex, as described in WO 90/10448. It is understood that the use of antisense molecules or knock out and knock in models may also be used in screening assays as discussed above, in addition to methods of treatment.

Thus, in one embodiment, methods of modulating angiogenesis in cells or organisms are provided. In one embodiment, the methods comprise administering to a cell an anti-angiogenesis antibody that reduces or eliminates the biological activity of an endogeneous angiogenesis protein. Alternatively, the methods comprise administering to a cell or organism a recombinant nucleic acid encoding an angiogenesis protein. This may be accomplished in any number of ways. In a preferred embodiment, for example when the angiogenesis sequence is down-regulated in angiogenesis, such state may be reversed by increasing the amount of angiogenesis gene product in the cell. This can be accomplished, e.g., by overexpressing the endogeneous angiogenesis gene or administering a gene encoding the angiogenesis sequence, using known gene-therapy techniques, for example. In a preferred embodiment, the gene therapy techniques include the incorporation of the exogenous gene using enhanced homologous recombination (EHR), for example as described in PCT/US93/03868, hereby incorporated by reference in its entireity. Alternatively, for example when the angiogenesis sequence is up-regulated in angiogenesis, the activity of the endogeneous angiogenesis gene is decreased, for example by the administration of a angiogenesis antisense nucleic acid or other inhibitor, such as RNAi.

In one embodiment, the angiogenesis eproteins of the present invention may be used to generate polyclonal and monoclonal antibodies to angiogenesis proteins. Similarly, the angiogenesis proteins can be coupled, using standard technology, to affinity chromatography columns. These columns may then be used to purify angiogenesis antibodies useful for production, diagnostic, or therapeutic purposes. In a preferred embodiment, the antibodies are generated to epitopes unique to a angiogenesis protein; that is, the antibodies show little or no cross-reactivity to other proteins. The angiogenesis antibodies may be coupled to standard affinity chromatography columns and used to purify angiogenesis proteins. The antibodies may also be used as blocking polypeptides, as outlined above, since they will specifically bind to the angiogenesis protein.

Methods of identifying variant angiogenesis-associated sequences

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Without being bound by theory, expression of various angiogenesis sequences is correlated with angiogenesis. Accordingly, disorders based on mutant or variant angiogenesis genes may be determined. In one embodiment, the invention provides methods for identifying cells containing variant angiogenesis genes, e.g., determining all or part of the sequence of at least one endogeneous angiogenesis genes in a cell. This may be accomplished using any number of sequencing techniques. In a preferred embodiment, the invention provides methods of identifying the angiogenesis genotype of an individual, e.g., determining all or part of the sequence of at least one angiogenesis gene of the individual. This is generally done in at least one tissue of the individual, and may include the evaluation of a number of tissues or different samples of the same tissue. The method may include comparing the sequence of the sequenced angiogenesis gene to a known angiogenesis gene, i.e., a wild-type gene.

The sequence of all or part of the angiogenesis gene can then be compared to the sequence of a known angiogenesis gene to determine if any differences exist. This can be done using any number of known homology programs, such as Bestfit, etc. In a preferred embodiment, the presence of a a difference in the sequence between the angiogenesis gene of the patient and the known angiogenesis gene correlates with a disease state or a propensity for a disease state, as outlined herein.

In a preferred embodiment, the angiogenesis genes are used as probes to determine the number of copies of the angiogenesis gene in the genome.

In another preferred embodiment, the angiogenesis genes are used as probes to determine the chromosomal localization of the angiogenesis genes. Information such as

chromosomal localization finds use in providing a diagnosis or prognosis in particular when chromosomal abnormalities such as translocations, and the like are identified in the angiogenesis gene locus.

5 Administration of pharmaceutical and vaccine compositions

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In one embodiment, a therapeutically effective dose of an angiogenesis protein or modulator thereof, is administered to a patient. By "therapeutically effective dose" herein is meant a dose that produces effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques (e.g., Ansel et al., Pharmaceuitcal Dosage Forms and Drug Delivery, Lippincott, Williams & Wilkins Publishers, ISBN:0683305727; Lieberman (1992) Pharmaceutical Dosage Forms (vols. 1-3), Dekker, ISBN 0824770846, 082476918X, 0824712692, 0824716981; Lloyd (1999) The Art, Science and Technology of Pharmaceutical Compounding, Amer. Pharmacutical Assn, ISBN 0917330889; and Pickar (1999) Dosage Calculations, Delmar Pub, ISBN 0766805042). As is known in the art, adjustments for angiogenesis degradation, systemic versus localized delivery, and rate of new protease synthesis, as well as the age, body weight, general health, sex, diet, time of administration, drug interaction and the severity of the condition may be necessary, and will be ascertainable with routine experimentation by those skilled in the art.

A "patient" for the purposes of the present invention includes both humans and other animals, particularly mammals. Thus the methods are applicable to both human therapy and veterinary applications. In the preferred embodiment the patient is a mammal, preferably a primate, and in the most preferred embodiment the patient is human.

The administration of the angiogenesis proteins and modulators thereof of the present invention can be done in a variety of ways as discussed above, including, but not limited to, orally, subcutaneously, intravenously, intranasally, transdermally, intraperitoneally, intramuscularly, intrapulmonary, vaginally, rectally, or intraocularly. In some instances, for example, in the treatment of wounds and inflammation, the angiogenesis proteins and modulators may be directly applied as a solution or spray.

The pharmaceutical compositions of the present invention comprise an angiogenesis protein in a form suitable for administration to a patient. In the preferred embodiment, the pharmaceutical compositions are in a water soluble form, such as being present as pharmaceutically acceptable salts, which is meant to include both acid and base addition salts. "Pharmaceutically acceptable acid addition salt" refers to those salts that retain

the biological effectiveness of the free bases and that are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid and the like, and organic acids such as acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like. "Pharmaceutically acceptable base addition salts" include those derived from inorganic bases such as sodium, potassium, lithium, ammonium, calcium, magnesium, iron, zinc, copper, manganese, aluminum salts and the like. Particularly preferred are the ammonium, potassium, sodium, calcium, and magnesium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, substituted amines including naturally occurring substituted amines, cyclic amines and basic ion exchange resins, such as isopropylamine, trimethylamine, diethylamine, triethylamine, tripropylamine, and ethanolamine.

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The pharmaceutical compositions may also include one or more of the following: carrier proteins such as serum albumin; buffers; fillers such as microcrystalline cellulose, lactose, corn and other starches; binding agents; sweeteners and other flavoring agents; coloring agents; and polyethylene glycol.

The pharmaceutical compositions can be administered in a variety of unit dosage forms depending upon the method of administration. For example, unit dosage forms suitable for oral administration include, but are not limited to, powder, tablets, pills, capsules and lozenges. It is recognized that angiogenesis protein modulators (e.g., antibodies, antisense constructs, ribozymes, small organic molecules, etc.) when administered orally, should be protected from digestion. This is typically accomplished either by complexing the molecule(s) with a composition to render it resistant to acidic and enzymatic hydrolysis, or by packaging the molecule(s) in an appropriately resistant carrier, such as a liposome or a protection barrier. Means of protecting agents from digestion are well known in the art.

The compositions for administration will commonly comprise an angiogenesis protein modulator dissolved in a pharmaceutically acceptable carrier, preferably an aqueous carrier. A variety of aqueous carriers can be used, e.g., buffered saline and the like. These solutions are sterile and generally free of undesirable matter. These compositions may be sterilized by conventional, well known sterilization techniques. The compositions may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions such as pH adjusting and buffering agents, toxicity adjusting agents

and the like, for example, sodium acetate, sodium chloride, potassium chloride, calcium chloride, sodium lactate and the like. The concentration of active agent in these formulations can vary widely, and will be selected primarily based on fluid volumes, viscosities, body weight and the like in accordance with the particular mode of administration selected and the patient's needs (e.g., Remington's Pharmaceutical Science, 15th ed., Mack Publishing Company, Easton, Pennsylvania (1980) and Goodman and Gillman, The Pharmacologial Basis of Therapeutics, (Hardman, J.G, Limbird, L.E, Molinoff, P.B., Ruddon, R.W, and Gilman, A.G., eds) TheMcGraw-Hill Companies, Inc., 1996).

Thus, a typical pharmaceutical composition for intravenous administration would be about 0.1 to 10 mg per patient per day. Dosages from 0.1 up to about 100 mg per patient per day may be used, particularly when the drug is administered to a secluded site and not into the blood stream, such as into a body cavity or into a lumen of an organ. Substantially higher dosages are possible in topical administration. Actual methods for preparing parenterally administrable compositions will be known or apparent to those skilled in the art, e.g., Remington's Pharmaceutical Science and Goodman and Gillman, The Pharmacologial Basis of Therapeutics, supra.

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The compositions containing modulators of angiogenesis proteins can be administered for therapeutic or prophylactic treatments. In therapeutic applications, compositions are administered to a patient suffering from a disease (e.g., a cancer) in an amount sufficient to cure or at least partially arrest the disease and its complications. An amount adequate to accomplish this is defined as a "therapeutically effective dose." Amounts effective for this use will depend upon the severity of the disease and the general state of the patient's health. Single or multiple administrations of the compositions may be administered depending on the dosage and frequency as required and tolerated by the patient. In any event, the composition should provide a sufficient quantity of the agents of this invention to effectively treat the patient. An amount of modulator that is capable of preventing or slowing the development of cancer in a mammal is referred to as a "prophylactically effective dose." The particular dose required for a prophylactic treatment will depend upon the medical condition and history of the mammal, the particular cancer being prevented, as well as other factors such as age, weight, gender, administration route, efficiency, etc. Such prophylactic treatments may be used, e.g., in a mammal who has previously had cancer to prevent a recurrence of the cancer, or in a mammal who is suspected of having a significant likelihood of developing cancer.

It will be appreciated that the present angiogenesis protein-modulating compounds can be administered alone or in combination with additional angiogenesis modulating compounds or with other therapeutic agent, e.g., other anti-cancer agents or treatments.

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In numerous embodiments, one or more nucleic acids, e.g., polynucleotides comprising nucleic acid sequences set forth in Tables 1-8, such as antisense polynucleotides or ribozymes, will be introduced into cells, in vitro or in vivo. The present invention provides methods, reagents, vectors, and cells useful for expression of angiogenesis-associated polypeptides and nucleic acids using in vitro (cell-free), ex vivo or in vivo (cell or organism-based) recombinant expression systems.

The particular procedure used to introduce the nucleic acids into a host cell for expression of a protein or nucleic acid is application specific. Many procedures for introducing foreign nucleotide sequences into host cells may be used. These include the use of calcium phosphate transfection, spheroplasts, electroporation, liposomes, microinjection, plasma vectors, viral vectors and any of the other well known methods for introducing cloned genomic DNA, cDNA, synthetic DNA or other foreign genetic material into a host cell (see, e.g., Berger and Kimmel, Guide to Molecular Cloning Techniques, Methods in Enzymology volume 152 Academic Press, Inc., San Diego, CA (Berger), F.M. Ausubel et al., eds., Current Protocols, a joint venture between Greene Publishing Associates, Inc. and John Wiley & Sons, Inc., (supplemented through 1999), and Sambrook et al., Molecular Cloning - A Laboratory Manual (2nd Ed.), Vol. 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989.

In a preferred embodiment, angiogenesis proteins and modulators are administered as therapeutic agents, and can be formulated as outlined above. Similarly, angiogenesis genes (including both the full-length sequence, partial sequences, or regulatory sequences of the angiogenesis coding regions) can be administered in a gene therapy application. These angiogenesis genes can include antisense applications, either as gene therapy (i.e. for incorporation into the genome) or as antisense compositions, as will be appreciated by those in the art.

Angiogenesis polypeptides and polynucleotides can also be administered as vaccine compositions to stimulate HTL, CTL and antibody responses.. Such vaccine compositions can include, for example, lipidated peptides (e.g., Vitiello, A. et al., J. Clin. Invest. 95:341, 1995), peptide compositions encapsulated in poly(DL-lactide-co-glycolide) ("PLG") microspheres (see, e.g., Eldridge, et al., Molec. Immunol. 28:287-294, 1991: Alonso

et al., Vaccine 12:299-306, 1994; Jones et al., Vaccine 13:675-681, 1995), peptide compositions contained in immune stimulating complexes (ISCOMS) (see, e.g., Takahashi et al., Nature 344:873-875, 1990; Hu et al., Clin Exp Immunol. 113:235-243, 1998), multiple antigen peptide systems (MAPs) (see e.g., Tam, J. P., Proc. Natl. Acad. Sci. U.S.A. 85:5409-5413, 1988; Tam, J.P., J. Immunol. Methods 196:17-32, 1996), peptides formulated as 5 multivalent peptides; peptides for use in ballistic delivery systems, typically crystallized peptides, viral delivery vectors (Perkus, M. E. et al., In: Concepts in vaccine development, Kaufmann, S. H. E., ed., p. 379, 1996; Chakrabarti, S. et al., Nature 320:535, 1986; Hu, S. L. et al., Nature 320:537, 1986; Kieny, M.-P. et al., AIDS Bio/Technology 4:790, 1986; Top, F. 10 H. et al., J. Infect. Dis. 124:148, 1971; Chanda, P. K. et al., Virology 175:535, 1990), particles of viral or synthetic origin (e.g., Kofler, N. et al., J. Immunol. Methods. 192:25, 1996; Eldridge, J. H. et al., Sem. Hematol. 30:16, 1993; Falo, L. D., Jr. et al., Nature Med. 7:649, 1995), adjuvants (Warren, H. S., Vogel, F. R., and Chedid, L. A. Annu. Rev. Immunol. 4:369, 1986; Gupta, R. K. et al., Vaccine 11:293, 1993), liposomes (Reddy, R. et al., J. Immunol. 148:1585, 1992; Rock, K. L., Immunol. Today 17:131, 1996), or, naked or particle 15 absorbed cDNA (Ulmer, J. B. et al., Science 259:1745, 1993; Robinson, H. L., Hunt, L. A., and Webster, R. G., Vaccine 11:957, 1993; Shiver, J. W. et al., In: Concepts in vaccine development, Kaufmann, S. H. E., ed., p. 423, 1996; Cease, K. B., and Berzofsky, J. A., Annu. Rev. Immunol. 12:923, 1994 and Eldridge, J. H. et al., Sem. Hematol. 30:16, 1993). 20 Toxin-targeted delivery technologies, also known as receptor mediated targeting, such as those of Avant Immunotherapeutics, Inc. (Needham, Massachusetts) may also be used.

Vaccine compositions often include adjuvants. Many adjuvants contain a substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A, Bortadella pertussis or Mycobacterium tuberculosis derived proteins. Certain adjuvants are commercially available as, for example, Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI); Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ); AS-2 (SmithKline Beecham, Philadelphia, PA); aluminum salts such as aluminum hydroxide gel (alum) or aluminum phosphate; salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars; cationically or anionically derivatized polysaccharides; polyphosphazenes; biodegradable microspheres; monophosphoryl lipid A and quil A. Cytokines, such as GM-CSF, interleukin-2, -7, -12, and other like growth factors, may also be used as adjuvants.

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Vaccines can be administered as nucleic acid compositions wherein DNA or RNA encoding one or more of the polypeptides, or a fragment thereof, is administered to a patient. This approach is described, for instance, in Wolff et. al., Science 247:1465 (1990) as well as U.S. Patent Nos. 5,580,859; 5,589,466; 5,804,566; 5,739,118; 5,736,524; 5,679,647; WO 98/04720; and in more detail below. Examples of DNA-based delivery technologies include "naked DNA", facilitated (bupivicaine, polymers, peptide-mediated) delivery, cationic lipid complexes, and particle-mediated ("gene gun") or pressure-mediated delivery (see, e.g., U.S. Patent No. 5,922,687).

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For therapeutic or prophylactic immunization purposes, the peptides of the invention can be expressed by viral or bacterial vectors. Examples of expression vectors include attenuated viral hosts, such as vaccinia or fowlpox. This approach involves the use of vaccinia virus, for example, as a vector to express nucleotide sequences that encode angiogenic polypeptides or polypeptide fragments. Upon introduction into a host, the recombinant vaccinia virus expresses the immunogenic peptide, and thereby elicits an immune response. Vaccinia vectors and methods useful in immunization protocols are described in, e.g., U.S. Patent No. 4,722,848. Another vector is BCG (Bacille Calmette Guerin). BCG vectors are described in Stover et al., Nature 351:456-460 (1991). A wide variety of other vectors useful for therapeutic administration or immunization e.g. adeno and adeno-associated virus vectors, retroviral vectors, Salmonella typhi vectors, detoxified anthrax toxin vectors, and the like, will be apparent to those skilled in the art from the description herein (see, e.g., Shata et al. (2000) Mol Med Today, 6: 66-71; Shedlock et al., J Leukoc Biol 68,:793-806, 2000; Hipp et al., In Vivo 14:571-85, 2000).

Methods for the use of genes as DNA vaccines are well known, and include placing an angiogenesis gene or portion of an angiogenesis gene under the control of a regulatable promoter or a tissue-specific promoter for expression in an angiogenesis patient. The angiogenesis gene used for DNA vaccines can encode full-length angiogenesis proteins, but more preferably encodes portions of the angiogenesis proteins including peptides derived from the angiogenesis protein. In one embodiment, a patient is immunized with a DNA vaccine comprising a plurality of nucleotide sequences derived from an angiogenesis gene. For example, angiogenesis-associated genes or sequence encoding subfragments of an 30 angiogenesis protein are introduced into expression vectors and tested for their immunogenicity in the context of Class I MHC and an ability to generate cytotoxic T cell responses. This procedure provides for production of cytotoxic T cell responses against cells which present antigen, including intracellular epitopes.

In a preferred embodiment, the DNA vaccines include a gene encoding an adjuvant molecule with the DNA vaccine. Such adjuvant molecules include cytokines that increase the immunogenic response to the angiogenesis polypeptide encoded by the DNA vaccine. Additional or alternative adjuvants are available.

In another preferred embodiment angiogenesis genes find use in generating animal models of angiogenesis. When the angiogenesis gene identified is repressed or diminished in angiogenesic tissue, gene therapy technology, e.g., wherein antisense RNA directed to the angiogenesis gene will also diminish or repress expression of the gene. Animal models of angiogenesis find use in screening for modulators of an angiogenesis-associated sequence or modulators of angiogenesis. Similarly, transgenic animal technology including gene knockout technology, for example as a result of homologous recombination with an appropriate gene targeting vector, will result in the absence or increased expression of the angiogenesis protein. When desired, tissue-specific expression or knockout of the angiogenesis protein may be necessary.

It is also possible that the angiogenesis protein is overexpressed in angiogenesis. As such, transgenic animals can be generated that overexpress the angiogenesis protein. Depending on the desired expression level, promoters of various strengths can be employed to express the transgene. Also, the number of copies of the integrated transgene can be determined and compared for a determination of the expression level of the transgene. Animals generated by such methods find use as animal models of angiogenesis and are additionally useful in screening for modulators to treat angiogenesis or to evaluate a therapeutic entity.

Kits for Use in Diagnostic and/or Prognostic Applications

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For use in diagnostic, research, and therapeutic applications suggested above, kits are also provided by the invention. In the diagnostic and research applications such kits may include any or all of the following: assay reagents, buffers, angiogenesis-specific nucleic acids or antibodies, hybridization probes and/or primers, antisense polynucleotides, ribozymes, dominant negative angiogenesis polypeptides or polynucleotides, small molecules inhibitors of angiogenesis-associated sequences *etc.* A therapeutic product may include sterile saline or another pharmaceutically acceptable emulsion and suspension base.

In addition, the kits may include instructional materials containing directions (i.e., protocols) for the practice of the methods of this invention. While the instructional materials typically comprise written or printed materials they are not limited to such. Any

medium capable of storing such instructions and communicating them to an end user is contemplated by this invention. Such media include, but are not limited to electronic storage media (e.g., magnetic discs, tapes, cartridges, chips), optical media (e.g., CD ROM), and the like. Such media may include addresses to internet sites that provide such instructional materials.

The present invention also provides for kits for screening for modulators of angiogenesis-associated sequences. Such kits can be prepared from readily available materials and reagents. For example, such kits can comprise one or more of the following materials: an angiogenesis-associated polypeptide or polynucleotide, reaction tubes, and instructions for testing angiogenic-associated activity. Optionally, the kit contains biologically active angiogenesis protein. A wide variety of kits and components can be prepared according to the present invention, depending upon the intended user of the kit and the particular needs of the user. Diagnosis would typically involve evaluation of a plurality of genes or products. The genes will be selected based on correlations with important parameters in disease which may be identified in historical or outcome data.

It is understood that the examples described above in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All publications, sequences of accession numbers, and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference.

EXAMPLES

25 Example 1: Tissue Preparation, Labeling Chips, and Fingerprints

Purify total RNA from tissue using TRIzol Reagent

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Homogenize tissue samples in 1ml of TRIzol per 50mg of tissue using a Polytron 3100 homogenizer. The generator/probe used depends upon the tissue size. A generator that is too large for the amount of tissue to be homogenized will cause a loss of sample and lower RNA yield. TRIzol is added directly to frozen tissue, which is then homogenize. Following homogenization, insoluble material is removed by centrifugation at 7500 x g for 15 min in a Sorvall superspeed or 12,000 x g for 10 min. in an Eppendorf centrifuge at 4°C. The clear homogenate is transferred to a new tube for use. The samples may be frozen now at -60° to -70°C (and kept for at least one month). The homogenate is

mixed with 0.2ml of chloroform per 1ml of TRIzol reagent used in the original homogenization and incubated at room temp. for 2-3 minutes. The aqueous phase is then separated by centrifugation and transferred to a fresh tube and the RNA precipitated using isopropyl alcohol. The pellet is isolated by centrifugation, washed, air-dried, resuspended in an appropriate volume of DEPC H₂O, and the absorbance measured.

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Purification of poly A+ mRNA from total RNA is performed as follows. Heat an oligotex suspension to 37°C and mixing immediately before adding to RNA. The Elution Buffer is heated at 70°C. Warm up 2 x Binding Buffer at 65°C if there is precipitate in the buffer. Mix total RNA with DEPC-treated water, 2 x Binding Buffer, and Oligotex according to Table 2 on page 16 of the Oligotex Handbook. Incubate for 3 minutes at 65°C. Incubate for 10 minutes at room temperature. Centrifuge for 2 minutes at 14,000 to 18,000 g. Remove supernatant without disturbing Oligotex pellet. A little bit of solution can be left behind to reduce the loss of Oligotex. Gently resuspend in Wash Buffer OW2 and pipet onto spin column. Centrifuge the spin column at full speed for 1 minute. Transfer spin column to a new collection tube and gently resuspend in Wash Buffer OW2 and centrifuge as describe herein. Transfer spin column to a new tube and elute with 20 to 100 ul of preheated (70oC) Elution Buffer. Gently resuspend Oligotex resin by pipetting up and down. Centrifuge as above. Repeat elution with fresh elution buffer or use first eluate to keep the elution volume low. Read absorbance, using diluted Elution Buffer as the blank. Before proceeding with cDNA synthesis, precipitate the mRNA as follows: add 0.4 vol. of 7.5 M NH4OAc + 2.5 vol. of cold 100% ethanol. Precipitate at -20oC 1 hour to overnight (or 20-30 min. at -70oC). Centrifuge at 14,000-16,000 x g for 30 minutes at 4oC. Wash pellet with 0.5ml of 80%ethanol (-20oC) then centrifuge at 14,000-16,000 x g for 5 minutes at room temperature. Repeat 80% ethanol wash. Air dry the ethanol from the pellet in the hood.. Suspend pellet in DEPC H₂0 at lug/ul concentration.

To further Clean up total RNA using Qiagen's RNeasy kit, add no more than 100ug to an RNeasy column. Adjust sample to a volume of 100ul with RNase-free water. Add 350ul Buffer RLT then 250ul ethanol (100%) to the sample. Mix by pipetting (do not centrifuge) then apply sample to an RNeasy mini spin column. Centrifuge for 15 sec at >10,000rpm. Transfer column to a new 2-ml collection tube. Add 500ul Buffer RPE and centrifuge for 15 sec at >10,000rpm. Discard flowthrough. Add 500ul Buffer RPE and centrifuge for 15 sec at >10,000rpm. Discard flowthrough then centrifuge for 2 min at maximum speed to dry column membrane. Transfer column to a new 1.5-ml collection tube

and apply 30-50ul of RNase-free water directly onto column membrane. Centrifuge 1 min at >10,000rpm. Repeat elution. and read absorbance.

cDNA synthesis using Gibco's "SuperScript Choice System for cDNA Synthesis" kit

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First Strand cDNA synthesis is performed as follows. Use 5ug of total RNA or 1ug of polyA+ mRNA as starting material. For total RNA, use 2ul of SuperScript RT. For polyA+ mRNA, use 1ul of SuperScript RT. Final volume of first strand synthesis mix is 20ul. RNA must be in a volume no greater than 10ul. Incubate RNA with 1ul of 100pmol T7-T24 oligo for 10 min at 70C. On ice, add 7 ul of: 4ul 5X 1st Strand Buffer, 2ul of 0.1M DTT, and 1 ul of 10mM dNTP mix. Incubate at 37C for 2 min then add SuperScript RT. Incubate at 37C for 1 hour.

For the second strand synthesis, place 1st strand reactions on ice and add: 91ul DEPC H₂0; 30ul 5X 2nd Strand Buffer; 3ul 10mM dNTP mix; 1ul 10U/ul E.coli DNA Ligase; 4ul 10U/ul E.coli DNA Polymerase; and 1ul 2U/ul RNase H. Mix and incubate 2 hours at 16C. Add 2ul T4 DNA Polymerase. Incubate 5 min at 16C. Add 10ul of 0.5M EDTA. A further clean-up of DNA is performed using phenol:chloroform:isoamyl Alcohol (25:24:1) purification.

In vitro Transcription (IVT) and labeling with biotin is performed as follows: Pipet 1.5ul of cDNA into a thin-wall PCR tube. Make NTP labeling mix by combining 2ul T7 10xATP (75mM) (Ambion); 2ul T7 10xGTP (75mM) (Ambion); 1.5ul T7 10xCTP (75mM) (Ambion); 3.75ul 10mM Bio-11-UTP (Boehringer-Mannheim/Roche or Enzo); 3.75ul 10mM Bio-16-CTP (Enzo); 2ul 10x T7 transcription buffer (Ambion); and 2ul 10x T7 enzyme mix (Ambion). The final volume is 20ul. Incubate 6 hours at 37°C in a PCR machine. The RNA can be furthered cleaned.

Fragmentation is performed as follows. 15 ug of labeled RNA is usually fragmented. Try to minimize the fragmentation reaction volume; a 10 ul volume is recommended but 20 ul is all right. Do not go higher than 20 ul because the magnesium in the fragmentation buffer contributes to precipitation in the hybridization buffer. Fragment RNA by incubation at 94 C for 35 minutes in 1 x Fragmentation buffer (5 x Fragmentation buffer is 200 mM Tris-acetate, pH 8.1; 500 mM KOAc; 150 mM MgOAc). The labeled RNA transcript can be analyzed before and after fragmentation. Samples can be heated to 65°C for 15 minutes and electrophoresed on 1% agarose/TBE gels to get an approximate idea of the transcript size range

For hybridization, 200 ul (10ug cRNA) of a hybridization mix is put on the chip. If multiple hybridizations are to be done (such as cycling through a 5 chip set), then it is recommended that an initial hybridization mix of 300 ul or more be made. The hybridization mix is: fragment labeled RNA (50ng/ul final conc.); 50 pM 948-b control oligo; 1.5 pM BioB; 5 pM BioC; 25 pM BioD; 100 pM CRE; 0.1mg/ml herring sperm DNA; 0.5mg/ml acetylated BSA; and 300 ul with 1xMES hyb buffer.

Labeling is performed as follows: The hybridization reaction includes non-biotinylated IVT (purified by RNeasy columns); IVT antisense RNA 4 μg:μl; random Hexamers (1 μg/μl) 4 μl and water to 14 ul. The reaciton is incubated at 70°C, 10 min. Reverse transcriptionis performed in the following reaction: 5X First Strand (BRL) buffer, 6 μl; 0.1 M DTT, 3 μl; 50X dNTP mix, 0.6 μl; H₂O, 2.4 μl; Cy3 or Cy5 dUTP (1mM), 3 μl; SS RT II (BRL), 1 μl in a final volume of 16 μl. Add to hybridization reaction. Incubate 30 min., 42°C. Add 1 μl SSII and incubate another hour. Put on ice. 50X dNTP mix (25mM of cold dATP, dCTP, and dGTP, 10mM of dTTP: 25 μl each of 100mM dATP, dCTP, and dGTP; 10 μl of 100mM dTTP to 15 μl H2O. dNTPs from Pharmacia)

RNA degradation is performed as follows. Add 86 µl H2O, 1.5 µl 1M NaOH/2mM EDTA and incubate at 65°C, 10 min.. For U-Con 30, 500 µl TE/sample spin at 7000g for 10 min, save flow through for purification. For Qiagen purification, suspend u-con recovered material in 500µl buffer PB and proceed using Qiagen protocol. For DNAse digestion, add 1 ul of 1/100 dil of DNAse/30ul Rx and incubate at 37°C for 15 min. Incubate at 5 min 95°C to denature the DNAse/

For sample preparation, add Cot-1 DNA, 10 μl; 50X dNTPs, 1 μl; 20X SSC, 2.3 μl; Na pyro phosphate, 7.5 μl; 10mg/ml Herring sperm DNA; 1ul of 1/10 dilution to 21.8 final vol. Dry in speed vac. Resuspend in 15 μl H20. Add 0.38 μl 10% SDS. Heat 95°C, 2 min and slow cool at room temp. for 20 min. Put on slide and hybridize overnight at 64°C. Washing after the hybridization: 3X SSC/0.03% SDS: 2 min., 37.5 mls 20X SSC+0.75mls 10% SDS in 250mls H2O; 1X SSC: 5 min., 12.5 mls 20X SSC in 250mls H2O; 0.2X SSC: 5 min., 2.5 mls 20X SSC in 250mls H2O. Dry slides and scan at appropriate PMT's and channels.

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Example 2. A model of angiogenesis is used to determine expression in angiogenesis

In the model of angiogenesis used to determine expression of angiogenesisassociated sequences, human umbilical vein endothelial cells (HUVEC) were obtained, e.g.,

as passage 1 (p1) frozen cells from Cascade Biologics (Oregon) and grown in maintenance medium: Medium 199 (Life Technologies) supplemented with 20% pooled human serum, 100 mg/ml heparin and 75 mg/ml endothelial cell growth supplements (Sigma) and gentamicin (Life Technologies). An *in vitro* cell system model was used in which 2x10⁵

HUVECs were cultured in 0.5 ml 3 mgs/ml plasminogen-depleted fibrinogen (Calbiochem, San Diego, CA) that was polymerized by the addition of 1 unit of maintenance medium supplemented with 100 ng/ml VEGF and HGF and 10 ng/ml TGF-a (R&D Systems, Minneapolis,MN) added (growth medium). The growth medium was replaced every 2 days. Samples for RNA were collected, *e.g.*, at 0, 2, 6, 15, 24, 48, and 96 hours of culture. The fibrin clots were placed in Trizol (Life Technologies) and disrupted using a Tissuemizer. Thereafter standard procedures were used for extracting the RNA (*e.g.*, Example 1).

Angiogenesis associated sequences thus identified are shown in Tables 1-8. As indicated, some of the Accession numbers include expression sequence tags (ESTs). Thus, in one embodiment herein, genes within an expression profile, also termed expression profile genes, include ESTs and are not necessarily full length.

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TABLE 1:

Unique Eos probeset identifier number Accession number used for previous patent fillings Exemptar Accession number, Genbank accession number Unigene number Unigene gene title Pkey: Accession: ExAccn: UnigeneID: Unigene Title: 5

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w		

10					
10	Pkey	Accession	ExAcon	UnigenelD	UnigeneTitle
	134404	AB000450	AB000450	Hs.82771	vaccinia related kinase 2
		AB002380	AF180681	Hs.6582	Rho guanine exchange factor (GEF) 12
15		AB003103	AA130080	Hs.4295	proteasome (prosome, macropain) 26S subunit, non-ATPase, 12
		AB004884	N27852 BE094848	Hs.57553 Hs.15113	tousled-like kinase 2 homogenitsate 1,2-dioxygenase (homogenitsate oxidase)
		AF000573_ma1 AF008937	AF008937	Hs.102178	syntaxin 16
		AF009301	AB011169	Hs.20141	similar to S. cerevisiae SSM4
20		AF009368	AF029674	Hs.173422	KIAA1605 protein
		D00591	NM_001269	Hs.84746	chromosome condensation 1
		D00760 D11139	AA294921 AA469369	Hs.250811 Hs.5831	v-rat simian leukemia viral oncogene homolog B (ras related; GTP binding protein) tissue Inhibitor of metalloproteinase 1 (erythroid potentiating activity, collagenase Inhibitor)
		D14657	H60720	Hs.81892	KIAA0101 gene product
25		D14878	AL037228	Hs.82043	D123 gene product
	101956	D17716	NM_002410	Hs.121502	mannosyl (alpha-1,6-)-glycoprotein beta-1,6-N-acetyl-glucosaminyltransferase
		D21090	M91401	Hs.178658	RAD23 (S. cerevisiae) homolog B
		D26135 D26528	NM_001346 D26528	Hs.89462 Hs.123058	diacy/glycerol kinase, gamma (90kD) DEAD/H (Asp-Gtu-Ala-Asp/His) box polypeptide 7 (RNA helicase, 52kD)
30		D30742	L24959	Hs.348	calcium/calmodulin-dependent protein kinase IV
		D31762	NM_012288	Hs.153954	TRAM-like protein
,		D31765	D31765	Hs.170114	KIAA0061 protein
		D31888	NM_015156	Hs.78398	KIAA0071 protein
35		D38128 D38500	D25418 D38500	Hs.393 Hs.278468	prostaglandin I2 (prostacyclin) receptor (IP) postmelotic segregation increased 2-like 4
55		D38551	N92036	Hs.81848	RAD21 (S. pombe) homolog
	100281	D42087	AF091035	Hs.184627	KIAA0118 protein
		D49396	AA331881	Hs.75454	peroxiredoxin 3
40		D55640 D63391	D55640 AW247529	Hs.6793	gb:Human monocyte PABL (pseudoautosomal boundary-like sequence) mRNA, done Mo2. platelet-activating factor acetylhydrolase, isoform lb, gamma subunit (29kD)
40		D63477	D63477	Hs.84087	KIAA0143 protein
	100338	D63483	D86864	Hs.57735	acetyl LDL receptor, SREC
		D64015	M96954	Hs.182741	TIA1 cytotoxic granule-associated RNA-binding protein-like 1
45		D79990	NM_014737	Hs.80905 Hs.184339	Ras association (RaIGDS/AF-6) domain family 2
45		D79997 D80010	NM_014791 BE613486	Hs.81412	KIAA0175 gene product lipin 1
		D84276	D84284	Hs.66052	CD38 antigen (p45)
		D86425	AW291587	Hs.82733	nidogen 2
50		D86978	D86978	Hs.84790	KIAA0225 protein
50		D87012 D87075	D87012 AF164142	Hs.194685 Hs.82042	topoisomerase (DNA) III beta solute carrier family 23 (nucleobase transporters), member 1
		D87432	D87432	Hs.10315	solute carrier family 7 (cationic amino acid transporter, y+ system), member 6
		D87448	AA013051	Hs.91417	topoisomerase (DNA) II binding protein
<i></i>		D87845	NM_000437	Hs.234392	platelet-activating factor acetylhydrolase 2 (40kD)
55		HG1098-HT1098 HG2167-HT2237		Hs.121489 Hs.301946	cystatin D lysosomal
		HG2415-HT2511		Hs.231444	Homo sapiens, Similar to hypothetical protein PRO1722, clone MGC:15692, mRNA, complete
	cds				
CO		HG2825-HT2949		Hs.142653	ret finger protein
60		HG2887-HT3031		A1368680	Hs.816 SRY (sex determining region Y)-box 2
	100899	HG4660-HT5073 HG4704-HT5146	1 12260	Hs.103042 Hs.172816	microtubule-associated protein 1B neuregulin 1
		HG884-HT884	AF002225	Hs.180686	ubiquitin protein ligase E3A (human papilloma virus E6-associated protein, Angelman syndrome)
	100950	HG919-HT919	AF128542	Hs.166846	polymerase (DNA directed), epsilon
65		J00212_f	J00212	11- 00000	Empirically selected from AFFX single probeset
	135407 130149		J04029 AW067805	Hs.99936 Hs.172665	keratin 10 (epidermolytic hyperkeratosis; keratosis palmaris et plantaris) methylenetetrahydrofolate dehydrogenase (NADP+ dependent), methenylletrahydrofolate
	131877	J04088	J04088	Hs.156346	topoisomerase (DNA) II alpha (170kD)
		J04543	J04543	Hs.78637	annexin A7
70	134786	L06139	T29618	Hs.89640	TEK tyrosine kinase, endothelial (venous malformations, multiple cutaneous and mucosal)
		L07540	AA460085 L08895	Hs.171075 Hs.78995	replication factor C (activator 1) 5 (36.5kD) MADS box transcription enhancer factor 2, polypeptide C (myocyte enhancer factor 2C)
		L08895 L11239	L11239	Hs.36993	gastrulation brain homeo box 1
		L11353	BE409525	Hs.902	neurofibrornin 2 (bilateral acoustic neuroma)
75	106432	L13773	AK000310	Hs.17138	hypothetical protein FLJ20303

	101152 L13800	Al984625	Hs.9884	spindle pole body protein
	135397 L14922	L14922	Hs.166563	replication factor C (activator 1) 1 (145kD)
	131687 L15189	BE297635	Hs.3069	heat shock 70kD protein 9B (mortalin-2)
_	101168 L15388	NM_005308		G protein-coupled receptor kinase 5
5	421155 L16895	H87879	Hs.102267	lysyl oxidase
	101226 L27476	AF083892	Hs.75608	tight junction protein 2 (zona occludens 2)
	133975 L27624	C18356	Hs.295944	tissue factor pathway inhibitor 2
	134739 L32976	NM_002419	Hs.89449	mitogen-activated protein kinase kinase kinase 11
10	130155 L33404	AA101043	Hs.151254	kallikrein 7 (chymotryptic, stratum comeum)
10	440538 L35263	W76332	Hs.79107	mitogen-activated protein kinase 14
	132813 L37347	BE313625	Hs.57435	solute carrier family 11 (proton-coupled divalent metal ion transporters), member 2
	101294 L40371	AF168418	Hs.116784	thyroid hormone receptor interactor 4
	101300 L40391	BE535511	Hs.74137	transmembrane trafficking protein
15	101310 L41607	L41607 AW250122	Hs.934 Hs.154879	glucosaminyl (N-acetyl) transferase 2, I-branching enzyme DiGeorge syndrome critical region gene DGSI; likely ortholog of mouse expressed sequence 2
13	130344 L77566 embryonic lethal	AVV250122	IIS. 1340/9	Dideotge syndicine dialeat region gene boot, axely outloing or mouse expressed sequence 2
	101381 M13928	AW675039	Hs.1227	aminolevulinate, delta-, dehydratase
	101668 M14016	AW005903	Hs.78601	uroporphyrinogen decarboxylase
	133780 M14219	AA557660	Hs.76152	decorin
20	101396 M15796	BE267931	Hs.78996	proliferating cell nuclear antigen
	101447 M21305	M21305		gb:Human alpha satellite and satellite 3 junction DNA sequence.
	101458 M22092	M22092		gb:Human neural cell adhesion molecule (N-CAM) gene, exon SEC and partial cds.
	101470 M22898	NM 000546	Hs.1846	tumor protein p53 (Li-Fraumeni syndrome)
	134604 M22995	NM_002884	Hs.865	RAP1A, member of RAS oncogene family
25	101478 M23379	NM_002890	Hs.758	RAS p21 protein activator (GTPase activating protein) 1
	406698 M24364	X03068	Hs.73931	major histocompatibility complex, class II, DQ beta 1
	133519 M24400	AW583062	Hs.74502	chymotrypsinogen B1
	131185 M25753	BE280074	Hs.23960	cyclin B1
20	134116 M27691	R84694	Hs.79194	cAMP responsive element binding protein 1
30	133999 M28213	AA535244	Hs.78305	RAB2, member RAS oncogene family
	130174 M29550	M29551	Hs.151531	protein phosphatase 3 (formerly 2B), catalytic subunit, beta isoform (calcineurin A beta)
	129963 M29971	M29971	Hs.1384	O-6-methylguanine-DNA methyltransferase
	132983 M30269	M30269	Hs.62041	nidogen (enactin) protein kinase, cAMP-dependent, regulatory, type II, beta
35	133900 M31158 101543 M31166	M31158 M31166	Hs.77439 _. Hs.2050	pentaxin-related gene, rapidly induced by IL-1 beta
55	101545 M31210		. Hs.154210	endothelial differentiation, sphingolipid G-protein-coupled receptor, 1
	101620 M55420	S55271	Hs.247930	Epsilon , IgE
	134691 M59979	AW382987	Hs.88474	prostaglandin-endoperoxide synthase 1 (prostaglandin G/H synthase and cyclooxygenase)
	133595 M62810	AA393273	Hs.75133	transcription factor 6-like 1 (mitochondrial transcription factor 1-like)
40	130425 M63838	AA243383	Hs.155530	interferon, gamma-inducible protein 16
	101700 M64710	D90337	Hs.247916	natriuretic peptide precursor C
	101714 M68874	M68874	Hs.211587	phospholipase A2, group IVA (cytosolic, calcium-dependent)
	134246 M74524	D28459	Hs.80512	ubiquitin-conjugating enzyme E2A (RAD6 homolog)
45	101760 M80254	M80254	Hs.173125	peptidylprolyl isomerase F (cyclophilin F)
45.	133948 M81780_cds3	X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid lysosomal (acid sphingomyelinase)
	101791 M83822	M83822	Hs.62354	cell division cycle 4-like DNA segment, numerous copies, expressed probes (GS1 gene)
	101812 M86934 101813 M87338	BE439894 NM_002914	Hs.78991 Hs.139226	replication factor C (activator 1) 2 (40kD)
	133396 M96326_ma1	M96326	Hs.72885	azurocidin 1 (cationic antimicrobial protein 37)
50	135152 M96954	M96954	Hs.182741	TIA1 cytotoxic granule-associated RNA-binding protein-like 1
50	129026 M98833	AL120297	Hs.108043	Friend leukemia virus integration 1
	101901 S66793	H38026	Hs.308	arrestin 3, retinal (X-arrestin)
	134831 S72370	AA853479	Hs.89890	рупиvate carboxylase
	134039 S78569	NM_002290	Hs.78672	laminin, alpha 4
55	134395 S79873	AA456539	Hs.8262	lysosomal
	101975 \$83325	AA079717	Hs.283664	aspartate beta-hydroxytase
	101977 S83364	AF112213	Hs.184062	putative Rab5-interacting protein
	101978 S83365	BE561610	Hs.5809	putative transmembrane protein; homolog of yeast Golgi membrane protein Yff1p (Ylp1p-
CO	interacting factor)			
60	101998 U01212	U01212	Hs.248153	olfactory marker protein
	102003 U01922	U01922	Hs.125565	translocase of Inner mitochondrial membrane 8 (yeast) homolog A
	102007 U02556	U02556	Hs.75307	t-complex-associated-testis-expressed 1-like
	102009 U02680	BE245149 U03272	Hs.82643	protein tyrosine kinase 9
65	416658 U03272 132951 U04209	AW821182	Hs.79432 Hs.61418	fibrillin 2 (congenital contractural aractmodactyly) microfibrillar-associated protein 1
05	135389 U05237	U05237	Hs.99872	fetal Alzheimer antigen
	102048 U07225	U07225	Hs.339	purinergic receptor P2Y, G-protein coupled, 2
	130145 U07620	U34820	Hs.151051	miliogen-activated protein kinase 10
	303153 U09759	U09759	Hs.246857	mitogen-activated protein kinase 9
70	420269 U09820	U72937	Hs.96264	alpha thalassemia/mental retardation syndrome X-linked (RAD54 (S. cerevisiae) homolog)
. •	102095 U11313	U11313	Hs.75760	sterol carrier protein 2
	102123 U14518	NM_001809	Hs.1594	centromere protein A (17kD)
	102126 U14575	AW950870	Hs.78961	protein phosphatase 1, regulatory (inhibitor) subunit 8
	102133 U15173	AU076845	Hs.155596	BCL2/adenovirus E1B 19kD-interacting protein 2
75	102139 U15932	NM_004419	Hs.2128	dual specificity phosphatase 5
	102162 U18291	AA450274	Hs.1592	CDC16 (cell division cycle 16, S. cerevisiae, homolog)

	102164 U18300	NIA 000107	Ua 77600	domese enceits DNA hinding protein 2 (4900)
	427653 U18383	NM_000107 AA159001	Hs.77602 Hs.180069	damage-specific DNA binding protein 2 (48kD) nuclear respiratory factor 1
	131817 U20536	U20536	Hs.3280	caspase 6, apoptosis-related cysteine protease
	102200 U21551	AA232362	Hs.157205	branched chain aminotransferase 1, cytosolic
5	102210 U23028	BE619413	Hs.2437	eukaryotic translation initiation factor 2B, subunit 5 (epsilon, 82kD)
	102214 U23752	U23752	Hs.32964	SRY (sex determining region Y)-box 11
	132811 U25435	U25435	Hs.57419	CCCTC-binding factor (zinc finger protein) stanniocalcin 1
	131319 U25997 102256 U28251_cds2	NM_003155 U28251	Hs.25590 Hs.53237	ESTS, Highly similar to Z169_HUMAN ZINC FINGER PROTEIN 169 [H.saplens]
10	132316 U28831	U28831	Hs.44566	KIAA1641 protein
	102269 U30245	U30245		gb:Human myelomonocytic specific protein (MNDA) gene, 5' flanking sequence and complete
	exon 1.			
	134365 U32315	AA568906	Hs.82240	syntaxin 3A
15	102293 U32439 102298 U32849	AF090116	Hs.79348	regulator of G-protein signalling 7
13	102325 U35139	AA382169 Al815867	Hs.54483 Hs.50130	N-myc (and STAT) interactor necdin (mouse) homolog
	302344 U36764	BE303044	Hs.192023	eukaryotic translation initiation factor 3, subunit 2 (beta, 36kD)
	102361 U39400	AA223616	Hs.75859	chromosome 11 open reading frame 4
20	102367 U39657	U39656	Hs.118825	mitogen-activated protein kinase kinase 6
20	102388 U41344	AA362907	Hs.76494	proline arginine-rich end leucine-rich repeat protein
	102394 U41766 129829 U41813	NM_003816 AF010258	Hs.2442 Hs.127428	a disintegrin and metalloproteinase domain 9 (maltrin gamma) homeo box A9
	102251 U41815	NM_004398	Hs.41706	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 10 (RNA helicase)
	102409 U43286	BE300330	Hs.118725	selenophosphate synthetase 2
25	133746 U44378	AW410035	Hs.75862	MAD (mothers against decapentaplegic, Drosophila) homolog 4
	102423 U44754	Z47542	Hs.179312	small nuclear RNA activating complex, polypeptide 1, 43kD
	132828 U47011_cds1	AB014615	Hs.57710	fibroblast growth factor 8 (androgen-induced)
	130441 U47077 102450 U48251	U63630 U48251	Hs.155637 Hs.75871	protein kinase, DNA-activated, catalytic polypeptide protein kinase C binding protein 1
30	129350 U50535	U50535	Hs.110630	Human BRCA2 region, mRNA sequence CG006
20	102534 U56833	U96759	Hs.198307	von Hippel-Lindau binding protein 1
	130457 U58091	AB014595	Hs.155976	cuttin 48
	135065 U58837	AA019401	Hs.93909	cyclic nucleotide gated channel beta 1
25	102560 U59289	R97457	Hs.63984	cadherin 13, H-cadherin (heart)
35	102567 U59863 134305 U67122	U63830 U61397	Hs.146847 Hs.81424	TRAF family member-associated NFKB activator ubiquitin-like 1 (sentrin)
	102638 U67319	U67319	Hs.9216	caspase 7, apoptosis-related cysteine protease
	132736 U68019	AW081883	Hs.288261	Homo sapiens cDNA: FLJ23037 fis, clone LNG02036, highly similar to HSU68019 Homo
	sapiens mad protein home	olog (hMAD-3) m	1RNA	
40	133070 U69611	U92649	Hs.64311	a disintegrin and metalloproteinase domain 17 (tumor necrosis factor, alpha, converting enzyme)
	102663 U70322	NM_002270	Hs.168075	karyopherin (importin) beta 2
	134660 U73524 102735 U79267	U73524 AF111106	Hs.87465 Hs.3382	ATP/GTP-binding protein protein phosphatase 4, regulatory subunit 1
	102741 U79291	AW959829	Hs.83572	hypothetical protein MGC14433
45	101175 U82671_cds2	U82671	Hs.36980	melanoma antigen, family A, 2
	132164 U84573	AI752235	Hs.41270	procollagen-lysine, 2-oxoglutarate 5-dioxygenase (lysine hydroxylase) 2
	102823 U90914	D85390	Hs.5057	carboxypeptidase D
	102826 U91316 102831 U91932	NM_007274 AA262170	Hs.8679 Hs.80917	cytosolic acyl coenzyme A thioester hydrolase adaptor-related protein complex 3, sigma 1 subunit
50	102846 U96131	BE264974	Hs.6566	thyroid hormone receptor interactor 13
	129777 U97018	U97018	Hs.12451	echinoderm microtubula-associated protein-like
	134161 U97188	AA634543	Hs.79440	IGF-II mRNA-binding protein 3
	134854 V00503	J03464	Hs.179573	collagen, type I, alpha 2
55	302363 X04327 133708 X06389	AW163799 AI018666	Hs.198365 Hs.75667	2,3-bisphosphoglycerate mutase synaptophysin
55	125701 X07496	T72104	Hs.93194	apolipoprotein A-l
	102915 X07820	X07820	Hs.2258	matrix metalloproteinase 10 (stromelysin 2)
	134656 X14787	AI750878	Hs.87409	thrombospondin 1
60	413858 X15525_ma1	NM_001610	Hs.75589	acid phosphatase 2, lysosomal
60	102968 X16396	AU076611	Hs.154672	methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate
	cyclohydrolase 102971 X16609	X16609	Hs.183805	ankyrin 1, erythrocytic
	134037 X53586_ma1	A1808780	Hs.227730	integrin, alpha 6
	103023 X53793	AW500470	Hs.117950	multifunctional polypeptide similar to SAICAR synthetase and AIR carboxylase
65	103037 X54936	BE018302	Hs.2894	placental growth factor, vascular endothelial growth factor-related protein
	130282 X55740	BE245380	Hs.153952	5' nucleotidase (CD73)
	134542 X57025 128568 X60673_ma1	M14156 H12912	Hs.85112 Hs.274691	insulin-like growth factor 1 (somatomedin C) adenytate kinase 3
	103093 X60708	S79876	Hs.44926	dipeptidy(peptidase IV (CD26, adenosine dearninase complexing protein 2)
70	133606 X62048	U10564	Hs.75188	wes1+ (S. pombe) homolog
		X63094	Hs.283822	Rhesus blood group, D antigen
	129063 X63097			
	424460 X63563	BE275979	Hs.296014	polymerase (RNA) II (DNA directed) polypeptide B (140kD)
	424460 X63563 133227 X64037	BE275979 AW977263	Hs.68257	general transcription factor IIF, polypeptide 1 (74kD subunit)
75	424460 X63563 133227 X64037 103181 X69636	BE275979 AW977263 X69636	Hs.68257 Hs.334731	general transcription factor IIF, polypeptide 1 (74kD subunit) Homo sapiens, done IMAGE:3448306, mRNA, partial ods
75	424460 X63563 133227 X64037	BE275979 AW977263	Hs.68257	general transcription factor IIF, polypeptide 1 (74kD subunit)

	103208 X72841	AW411340	Hs.31314	retinoblastoma-binding protein 7
	129698 X74987	BE242144	Hs.12013	ATP-binding cassette, sub-family E (OABP), member 1
	131486 X83107	F06972	Hs.27372	BMX non-receptor tyrosine kinase
_	130729 X84194	AI963747	Hs.18573	acylphosphatase 1, erythrocyte (common) type
5	103334 X85753	NM_001260	Hs.25283	cyclin-dependent kinase 8
	132645 X87870	A1654712	Hs.54424	hepatocyte nuclear factor 4, alpha
	135094 X89066	NM_003304	Hs.250687	transient receptor potential channel 1
	103352 X89398_cds2	H09366	Hs.78853	uracii-DNA giyoosylase
10	103353 X89399	X89399	Hs.119274	RAS p21 protein activator (GTPase activating protein) 3 (Ins(1,3,4,5)P4-binding protein)
10	132173 X89426	X89426	Hs.41716	endothelial cell-specific molecule 1
	103371 X91247	X91247	Hs.13046	thioredoxin reductase 1
	131584 X91648	AA598509	Hs.29117	purine-rich element binding protein A
	103376 X92098	AL036166	Hs.323378	coated vesicle membrane protein
15	103378 X92110	AL119690	Hs.153618	HCGVIII-1 protein .
15	128510 X94703	X94703	Hs.296371	RAB28, member RAS oncogene family
	103410 X96506	AA158294	Hs.334879	DR1-associated protein 1 (negative cofactor 2 alpha)
	133490 X97230_f		Hs.274601	killer cell immunoglobulin-like receptor, three domains, long cytoplasmic tall, 1
	103438 X98263	AW175781	Hs.152720	M-phase phosphoprotein 6
20	103440 X98296	X98296	Hs.77578	ubiquitin specific protease 9, X chromosome (Drosophila fat facets related)
20	103452 X99584	NM_006936	Hs.85119	SMT3 (suppressor of mif two 3, yeast) homolog 1
	133536 Y00264	W25797.comp		amytoid beta (A4) precursor protein (protease nexin-li, Alzheimer disease)
	135185 Y07566	AW404908	Hs.96038	Ric (Drosophila)-like, expressed in many tissues
	118523 Y07759	Y07759	Hs.170157	myosin VA (heavy polypeptide 12, myoxin)
25	134662 Y07827 132083 Y07867	NM_007048	Hs.284283	butyrophilin, subfamily 3, member A1
23		BE386490	Hs.279663	Pirin
	103500 Y09443	AW408009	Hs.22580	alkylglycerone phosphate synthase
	134389 Y09858	Y09858	Hs.82577	spindlin-like
	132084 Y12394 103540 Z11559	NM_002267 NM_002197	Hs.3886 Hs.154721	karyopherin alpha 3 (importin alpha 4) aconitase 1, soluble
30	133152 Z11695	Z11695	Hs.324473	mitogen-activated protein kinase 1
50	103548 Z15005	Z15005	Hs.75573	centromere protein E (312kD)
	103612 Z46261	BE336654	Hs.70937	H3 histone family, member A
•	129092 AA011243_s	D56365	Hs.63525	poly(rC)-binding protein 2
	103692 AA018418	AW137912	Hs.227583	Homo sapiens chromosome X map Xp11.23 L-type calcium channel alpha-1 subunit
35	(CACNA1F) gene, complet	e cds: HSP27 p		riplete sequence; and JM1 protein, JM2 protein, and Hb2E genes, complete cds
J	103695 AA018758	AW207152	Hs.186600	ESTs
	129796 AA018804	BE218319	Hs.5807	GTPase Rab14
	132258 AA031993	AA306325	Hs.4311	SUMO-1 activating enzyme subunit 2
•	132683 AA044217	BE264633	Hs.143638	WD repeat domain 4
40	131887 AA046548	W17064	Hs.332848	SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily e,
	member 1			
	103723 AA057447_s	BE274312	Hs.214783	Homo sapiens cONA FLJ14041 fis, clone HEMBA1005780
	453368 AA058376	MARGOOG	Hs.288178	Homo sapiens cDNA FLJ11968 fis, clone HEMBB1001133
	100000 / 0 10000 / 0	W20296	113.200 110	The state of the s
	133260 AA083572	AA403045	Hs.6906	Homo saplens cDNA: FLJ23197 fis, clone REC00917
45	133260 AA083572 103765 AA085696			
45	133260 AA083572 103765 AA085696 103766 AA088744	AA403045 AA085696 AI920783	Hs.6906 Hs.169600 Hs.191435	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs
45	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688	AA403045 AA085696	Hs.6906 Hs.169600	Homo sapiens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein
45	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284	AA403045 AA085696 AI920783 BE244667 AA393968	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein
	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700	AA403045 AA085696 AI920783 BE244667	Hs.6906 Hs.169600 Hs.191435 Hs.296155	Homo sapiens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein
45 50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans]	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans
	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5
	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Wealdy similar to T22363 hypothetical protein F47G9.4 - Caenomabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD)
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50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842	Hs.6906 Hs.169600 Hs.191435 Hs.286155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Wealdy similar to T22363 hypothetical protein F47G9.4 - Caenomabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein
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50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation Initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein furnarate hydratasse similar to yeast Upf3, variant B
50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419	Hs.6506 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.103832	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratasse similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein)
50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA130016 119159 AA149507 129863 AA151005	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.192619 Hs.103832 Hs.103832 Hs.15020 Hs.129872	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein tumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9
50 55	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA19507 129863 AA151005 103850 AA187101	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101	Hs.6906 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55582 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenomabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895
50	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363	Hs.6906 Hs.199600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.20267	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein tumarate hydralase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330
50 55	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103855 AA195179_s 322026 AA203138	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.2302267 Hs.223675	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein
50 55	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.230267 Hs.23675 Hs.23675 Hs.278626	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratasse similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2
50 55	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55582 Hs.30670 Hs.192619 Hs.75553 Hs.103832 Hs.15020 Hs.129872 Hs.23194 Hs.23375 Hs.23875 Hs.23875 Hs.23875 Hs.23875 Hs.23875 Hs.23875	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratasse similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/AbI-Interacting protein ArgBP2 hypothetical protein FLJ12783
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA094800 119159 AA195179 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.230267 Hs.23675 Hs.23675 Hs.278626	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2
50 55	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA157101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans]	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA162922 AA206236 AI769067	Hs.6906 Hs.199600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.103832 Hs.123872 Hs.213194 Hs.302267 Hs.223675 Hs.278626 Hs.4944 Hs.127824	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydralase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268	Hs.6906 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.278626 Hs.4944 Hs.127824	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein tumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ123020 fis, clone LNG00943
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.2302267 Hs.233675 Hs.278626 Hs.4944 Hs.127824 Hs.6127 Hs.6127 Hs.6127 Hs.6127 Hs.6127 Hs.47438	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydralase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611 AF043117	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.23675 Hs.23675 Hs.278626 Hs.287575 Hs.278626 Hs.4944 Hs.127824	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydralasse similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E4B (homologous to yeast UFD2)
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA092968 132471 AA114885 134319 AA129547 103807 AA133016 119159 AA133016 119159 AA13016 119159 AA19517 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199	AA403045 AA085696 AI920783 BE2444667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611 AF043117 D42039	Hs.6906 Hs.19600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55582 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.302267 Hs.228675 Hs.228675 Hs.278626 Hs.4944 Hs.6127 Hs.6127 Hs.47438 Hs.24594 Hs.478871	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quakling QKI (KH domain RNA binding protein) spem associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E48 (homologous to yeast UFD2) mesoderm development candidate 2
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA167101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199 129013 AA313990	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA275268 AA249611 AF043117 D42039 AA371156	Hs.6906 Hs.19600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.230267 Hs.238675 Hs.278626 Hs.4944 Hs.127824 Hs.6127 Hs.47438 Hs.47438 Hs.47438 Hs.278671 Hs.17942	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaklng QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E4B (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA151005 103850 AA157101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 103861 AA206236 103861 AA206236 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199 129013 AA313990 129435 AA314256	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611 AF043117 D42039 AA371156 AF151852	Hs.6906 Hs.19600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55582 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.302267 Hs.228675 Hs.228675 Hs.278626 Hs.4944 Hs.6127 Hs.6127 Hs.47438 Hs.24594 Hs.478871	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein tumarate hydratase similar to yeast Upf3, variant B homolog of mouse quakling QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E4B (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA2026261 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199 129013 AA313990 129435 AA314256 103988 AA314389	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA275268 AA249611 AF043117 D42039 AA371156	Hs.6906 Hs.191435 Hs.296155 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.302267 Hs.23675 Hs.278626 Hs.4944 Hs.127824 Hs.6127 Hs.47438 Hs.24594 Hs.6127 Hs.47438 Hs.24594 Hs.78871 Hs.107942 Hs.111449	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaklng QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E4B (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein
5055606570	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199 129013 AA313990 129435 AA314256 103988 AA314389 104000 AA324364	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA29611 AF043117 D42039 AA371156 AF51852 AA314389	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.192872 Hs.129872 Hs.233675 Hs.278626 Hs.4944 Hs.127824 Hs.6127 Hs.47438 Hs.24594 Hs.6127 Hs.87438 Hs.24594 Hs.767942 Hs.107942 Hs.111449 Hs.42500	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation indicator 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E48 (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein GGI-94 protein ADP-ribosylation factor-like 5
50 55 60	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA149507 129863 AA151005 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA2026261 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA282640 134060 AA287199 129013 AA313990 129435 AA314256 103988 AA314389	AA403045 AA085696 AI920783 BE244667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611 AF043117 D42039 AA371156 AF151852 AA314389 AI146527	Hs.6906 Hs.169600 Hs.191435 Hs.296155 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55682 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.15020 Hs.129872 Hs.233675 Hs.233675 Hs.278626 Hs.4944 Hs.127824 Hs.6127 Hs.47438 Hs.24594 Hs.6127 Hs.107942 Hs.107942 Hs.111449 Hs.42500 Hs.80475	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation Initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydralase similar to yeast Upf3, variant B homolog of mouse quaking QKI (KH domain RNA binding protein) sperm associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E4B (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein CGI-94 protein ADP-rfbosykation factor-like 5 polymerase (RNA) II (DNA directed) polypeptide J (13.3kD)
5055606570	133260 AA083572 103765 AA085696 103766 AA088744 103767 AA089688 132051 AA091284 103773 AA092700 [C.elegans] 135289 AA092968 132729 AA094800 103794 AA100219 131471 AA114885 134319 AA129547 103807 AA133016 119159 AA094800 103850 AA187101 103855 AA195179_s 322026 AA203138 135300 AA203645 103861 AA206236 130634 AA227621 [C.elegans] 447735 AA248283 103909 AA249611 131236 AA248283 103909 AA249611 131236 AA26640 134060 AA267199 129013 AA313990 129435 AA314256 103988 AA314389 104000 AA324364 425284 AA329211_s	AA403045 AA085696 AI920783 BE2444667 AA393968 AI219323 AW372569 AW970843 AF244135 AA164842 BE304999 AW958264 AF142419 BE379765 AA187101 W02363 AW024973 AA142922 AA206236 AI769067 AA775268 AA249611 AF043117 D42039 AA371156 AF151852 AA314389 AI146527 AF155568	Hs.6906 Hs.19600 Hs.191435 Hs.296155 Hs.180145 Hs.101077 Hs.9788 Hs.55582 Hs.30670 Hs.192619 Hs.75653 Hs.103832 Hs.103832 Hs.103832 Hs.15020 Hs.129872 Hs.213194 Hs.302267 Hs.228575 Hs.278626 Hs.2444 Hs.127824 Hs.6127 Hs.827871 Hs.107942 Hs.111449 Hs.42500 Hs.42459 Hs.48475 Hs.155489	Homo saplens cDNA: FLJ23197 fis, clone REC00917 KIAA0826 protein ESTs CGI-100 protein HSPC030 protein ESTs, Weakly similar to T22363 hypothetical protein F47G9.4 - Caenorhabditis elegans hypothetical protein MGC10924 similar to Nedd4 WW-binding protein 5 eukaryotic translation initiation factor 3, subunit 7 (zeta, 66/67kD) hepatocellular carcinoma-associated antigen 66 KIAA1600 protein fumarate hydratases similar to yeast Upf3, variant B homolog of mouse quaklng QKI (KH domain RNA binding protein) spem associated antigen 9 hypothetical protein MGC10895 hypothetical protein FLJ10330 NPD009 protein Arg/Abl-Interacting protein ArgBP2 hypothetical protein FLJ12783 ESTs, Weakly similar to T28770 hypothetical protein W03D2.1 - Caenorhabditis elegans Homo sapiens cDNA: FLJ23020 fis, clone LNG00943 SH3 domain binding glutamic acid-rich protein ubiquitination factor E48 (homologous to yeast UFD2) mesoderm development candidate 2 DKFZP564M112 protein CGI-94 protein ADP-ribosylation factor-like 5 polymerase (RNA) II (DNA directed) polypeptide J (13.3kD) NS1-associated protein 1

	104104 AA422029	AA422029	Hs.143640	ESTs, Wealthy similar to hyperpolarization-activated cyclic nucleotide-gated channel hHCN2
	[H.sapiens]	NM_005754	Hs.220689	Ras-GTPase-activating protein SH3-domain-binding protein
	108154 AA425230 132091 AA447052	AW954243	Hs.170218	KIAA0251 protein
5	135073 AA452000	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (from clone DKFZp586E1624)
	131367 AA456687	AI750575	Hs.173933	nuclear factor I/A
	129593 AA487015_s	Al338247	Hs.98314	Homo sapiens mRNA; cDNA DKFZp586L0120 (from clone DKFZp586L0120)
	135266 AB002326	R41179	Hs.97393	KIAA0328 protein
10	133505 C01527	A1630124	Hs.324504 Hs.3838	Homo sapiens mRNA; cDNA DKFZp586J0720 (from clone DKFZp586J0720)
10	132064 C01714 134393 C01811_f	AA121098 W52642	Hs.8261	serum-indudible kinase hypothetical protein FLJ22393
	131427 C02352_s	AF151879	Hs.26706	CGI-121 protein
	133435 C02375	Al929357	Hs.323966	Homo sapiens clone H63 unknown mRNA
	104282 C14448	C14448	Hs.332338	EST
15	134827 D16611_s	BE314037	Hs.89866	coproporphyrinogen oxidase (coproporphyria, harderoporphyria)
	130443 D25216	D25216	Hs.155650	KIAA0014 gene product
	131742 D31352 132837 D58024_s	AA961420 AA370362	Hs.31433 Hs.57958	ESTs EGF-TM7-latrophilin-related protein
	130377 D80897	NM_014909	Hs.155182	KIAA1036 protein
20	104334 D82614	D82614	Hs.78771	phosphoglycerate kinase 1
	134593 D87845	NM_000437	Hs.234392	platelet-activating factor acetylhydrolase 2 (40kD)
	134731 D89377_i	D89377	Hs.89404	msh (Drosophila) homeo box homolog 2
	129913 H06583	NM_001310	Hs.13313	cAMP responsive element binding protein-like 2
25	131670 H40732	H03514 AA129551	Hs.10130 Hs.172129	ESTS
23	104394 H46617 104402 H56731	H56731	Hs.172125	Homo saplens cDNA: FLJ21409 fis, clone COL03924 ESTs
	129781 H75570	AA306090	Hs.124707	ESTs
	129077 H78886	N74724	Hs.108479	ESTs
	104417 H81241	Al819448	Hs.320861	Kruppel-like factor 8
30	134927 L36531	L36531	Hs.91296	integrin, alpha 8
	129280 M63154	M63154	Hs.110014	gastric intrinsic factor (vitamin B synthesis)
	134498 M63180 104460 M91504	AW246273 AW955705	Hs.84131 Hs.62604	threonyl-IRNA synthetase Homo sapiens, clone IMAGE:4299322, mRNA, partial cds
	104488 N56191	N56191	Hs.106511	protocadherin 17
35	131248 N78483	Al038989	Hs.332633	Bardel-Biedl syndrome 2
	129214 N79268	AL044335	Hs.109526	zinc finger protein 198
	130017 R14652	AK000096	Hs.143198	Inhibitor of growth family, member 3
	104530 R20459	AK001676 R22303	Hs.12457	hypothetical protein FLJ10814 gb:yh26b09.r1 Soares placenta Nb2HP Homo saptens cDNA clone IMAGE:130841 5', mRNA
40	104534 R22303 sequence.	R22303		gu.yizouda.i i dualea piazenia kuzhir honio sapiena cunia cidne iniadez. 130041 3, iniada.
10	104544 R33779	Al091173	Hs.222362	ESTs, Weakly similar to p40 [H.sapiens]
	133328 R36553	AW452738	Hs.265327	hypothetical protein DKFZp761I141
	104567 R64534	AA040620	Hs.5672	hypothetical protein AF140225
45	128562 R66475	AA923382	Hs.101490	ESTs
45	129575 R70621 130776 R79356	F08282	Hs.278428 Hs.19280	progestin induced protein cysteine-rich motor neuron 1
	104599 R84933	AF167706 AW815036	Hs.151251	ESTs
	104660 RC_AA007160		Hs.14846	Homo sapiens mRNA; cDNA DKFZp564D016 (from clone DKFZp564D016)
	104667 RC_AA007234		Hs.30098	ESTs
50	104718 RC_AA018409		Hs.36250	ESTs, Weakly similar to 138022 hypothetical protein [H.sapiens]
	104764 RC_AA025351		Hs.278585	ESTs
•	104786 RC_AA027168 104787 RC_AA027317		Hs.10031	KIAA0955 protein gb:ze97d11.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA done IMAGE:366933 3*
	similar to contains Alu re		mRNA seguen	
55	134079 RC_AA029423		Hs.171835	hypothetical protein FLJ 10889
	104804 RC_AA031357		Hs.31803	ESTs, Wealthy similar to N-WASP [H.sapiens]
	104865 RC_AA045136		Hs.22575	B-cell CLL/lymphoma 6, member B (zinc finger protein)
	130828 RC_AA053400		Hs.203213	ESTS
60	104907 RC_AA055829 WARNING ENTRY [H.sa		Hs.196701	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
OO	104943 RC_AA065217	AF072873	Hs.114218	frizzled (Drosophila) homolog 6
	105013 RC_AA116054	H63789	Hs.296288	ESTs, Wealdy similar to KIAA0638 protein [H.sapiens]
	105024 RC_AA126311		Hs.9879	ESTs
	132592 RC_AA129390		Hs.288850	Homo sapiens cDNA: FLJ22528 fis, done HRC12825
65	105038 RC_AA130273		Hs.9414	KIAA1488 protein
	105077 RC_AA142919 105096 RC_AA150205		Hs.234863 Hs.21599	Homo sapiens cDNA FLJ12082 fis, clone HEMB81002492 Kruppel-like factor 7 (ubiquitous)
	129215 RC_AA176867		Hs.126085	KIAA1497 protein
	105169 RC_AA180321		Hs.180789	S164 protein
70	132796 RC_AA180487		Hs.173159	transforming, acidic coiled-coil containing protein 1
	130401 RC_AA187634	BE396283	Hs.173987	eukaryofic translation initilation factor 3, subunit 1 (alpha, 35kD)
	105200 RC_AA195399		Hs.24641	cytoskeleton associated protein 2
	130114 RC_AA234717		Hs.14992	hypothetical protein FLJ11151 ESTs
75	105330 RC_AA234743 105337 RC_AA234957		Hs.22120 Hs.23200	myotubutarin related protein 1
, 5	129385 RC_AA235604		Hs.110950	Rag C protein
			•	

	105376 RC_AA236559	AW994032	Hs.8768	hypothetical protein FLJ 10849
	105397 RC_AA242868	AA814807	Hs.7395	hypothetical protein FLJ23182
	131962 RC_AA251776	AK000046	Hs.267448	hypothetical protein FLJ20039
	131991 RC_AA251909	AF053306	Hs.36708	budding uninhibited by benzimidazoles 1 (yeast homolog), beta
5	128658 RC_AA252672_s		Hs.324830	diptheria toxin resistance protein required for diphthamide biosynthesis (Saocharomyces)-like 2
•	105489 RC AA256157	AA256157	Hs.24115	Homo sapiens cDNA FLJ14178 fis, clone NT2RP2003339
	105508 RC_AA256680	AA173942	Hs.326416	Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
	105539 RC AA258873	AB040884	Hs.109694	KIAA1451 protein
	135172 RC_AA262727	AB028956	Hs.12144	KIAA1033 protein
10 .	131569 RC_AA281451	AL389951	Hs.271623	nucleoporin 50kD
10	132542 RC_AA281545	AL137751	Hs.263671	Homo sapiens mRNA; cDNA DKFZp434l0812 (from clone DKFZp434l0812); partial cds
	105643 RC AA282069	BE621719	Hs.173802	KIAA0603 gene product
	105659 RC_AA283044	AA283044	Hs.25625	hypothetical protein FLJ11323
	105666 RC_AA283930	AA426234	Hs.34906	ESTs, Weakly similar to T17210 hypothetical protein DKFZp434N041.1 [H.sapiens]
15	105674 RC_AA284755	AI609530	Hs.279789	histone deacetylase 3
13	105709 RC_AA291268	Al928962	Hs.26761	DKFZP586L0724 protein
	105722 RC_AA291927	Al922821	Hs.32433	ESTs
	105765 RC_AA343514	AA299688	Hs.24183	ESTs
	115951 RC_AA398109	BE546245	Hs.301048	sec13-like protein
20	105962 RC_AA405737	AW880358	Hs.339808	hypothetical protein FLJ10120
_ ,	105985 RC_AA406610	AA406610		gb:zv15b10.s1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:753691 3' similar to
	gb:X02067			3-2-13-13-13-13-13-13-13-13-13-13-13-13-13-
	106008 RC_AA411465	AB033888	Hs.8619	SRY (sex determining region Y)-box 18
	131216 RC_AA416886	AI815486	Hs.243901	Homo sapiens cDNA FLJ20738 fis, clone HEP08257
25	134222 RC_AA424013	AW855861	Hs.8025	Homo saplens clone 23767 and 23782 mRNA sequences
	113689 RC_AA424148	AB037850	Hs.16621	DKFZP434l116 protein
	106141 RC_AA424558	AF031463	Hs.9302	phosducin-like .
	130839 RC_AA424961_s		Hs.20141	similar to S. cerevisiae SSM4
	106157 RC_AA425367		Hs.34892	KIAA1323 protein
30	130777 RC_AA425921	AW135049	Hs.285418	Homo saplens cDNA FLJ10643 fis, clone NT2RP2005753, highly similar to Homo saplens I-1
	receptor			
	130561 RC_AA426220	AB011095	Hs.16032	KIAA0523 protein
	106196 RC_AA427735	AA525993	Hs.173699	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
	WARNING			
35	131878 RC_AA430673	AA083764	Hs.6101	hypothetical protein MGC3178
	133200 RC_AA432248	AB037715	Hs.183639	hypothetical protein FLJ10210
	106302 RC_AA435896	AA398859	Hs.18397	hypothetical protein FLJ23221
	106328 RC_AA436705	AL079559	Hs.28020	KIAA0766 gene product
	450534 RC_AA446561	Al570189	Hs.25132	KIAA0470 gene product
40	106423 RC_AA448238	AB020722	Hs.16714	Rho guanine exchange factor (GEF) 15
	133442 RC_AA448688	AL137663	Hs.7378	Homo sapiens mRNA; cDNA DKFZp434G227 (from done DKFZp434G227)
	439608 RC_AA449756	AW864696	Hs.301732	hypothetical protein MGC5306
	106477 RC_AA450303	R23324	Hs.41693	DnaJ (Hsp40) homolog, subfamily B, member 4
4.5	106503 RC_AA452411	AB033042	Hs.29679	cofactor required for Sp1 transcriptional activation, subunit 3 (130kD)
45	446999 RC_AA454566	AA151520	Hs.334822	hypothetical protein MGC4485
	106543 RC_AA454667	AA676939	Hs.69285	neuropilin 1
	130010 RC_AA456437	AA301116	Hs.142838	nucleolar phosphoprotein Nopp34
	106589 RC_AA456646	AK000933	Hs.28661	Homo sapiens cDNA FLJ10071 fis, clone HEMBA1001702
50	106593 RC_AA456826	AW296451	Hs.24605	ESTs
50	106596 RC_AA456981	AA452379	Hs.293552	ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE
	CONTAMINATION			Assessment to be a first to the description of the configuration of the
	134655 RC_AA458959	AF265208	Hs.123090	SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily f,
	member 1	*******	11- 000	M
55	106636 RC_AA459950	AW958037	Hs.286 ·	ribosomal protein L4
33	106654 RC_AA460449	AW075485	Hs.286049	phosphoserine aminotransferase
	131353 RC_AA463910	AW754182	11-00405	gb:RC2-CT0321-131199-011-001 CT0321 Homo sapiens cDNA, mRNA sequence
	106707 RC_AA464603	AK000566	Hs.98135	hypothetical protein FLJ20559
	131710 RC_AA464606	NM_015368	Hs.30985	pannexin 1 TIA1 cytotoxic granule-associated RNA-binding protein
60	106717 RC_AA465093	AA600357	Hs.239489	
00 .	131775 RC_AA465692 106747 RC_AA476473	AB014548 NM_007118	Hs.31921 Hs.171957	KIAA0648 protein triple functional domain (PTPRF interacting)
	106773 RC_AA478109	AA478109	Hs.188833	ESTS
	105781 RC_AA478474	AA330310	Hs.24181	ESTs
	106817 RC_AA480889	D61216	Hs.18672	ESTs
65	106846 RC_AA485223	AB037744	Hs.34892	KIAA1323 protein
03	106848 RC_AA485254	AA449014	Hs.121025	chromosome 11 open reading frame 5
	106856 RC_AA486183	W58353	Hs.285123	Homo sapiens mRNA full length insert cDNA done EUROIMAGE 2005779
	418699 RC_AA496936	BE539639	Hs.173030	ESTS, Weakly similar to ALUS_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION
	WARNING	2000003	1 10000	TOTO, TOTAL STIME IN PROOF INTERFERENCE OF THE PARTY OF OCCUPANT OF THE PARTY OF TH
70	107001 RC_AA598589	AJ926520	Hs.31016	putative DNA binding protein
, 5	130638 RC_AA598831_f		Hs.17121	ESTs
	107054 RC_AA600150	A1076459	Hs.15978	KIAA1272 protein
	107059 RC_AA608545	BE614410	Hs.23044	RAD51 (S. cerevisiae) homolog (E coli RecA homolog)
	107080 RC_AA609210	AL122043	Hs.19221	hypothetical protein DKFZp566G1424
75	107115 RC_AA610108	BE379623	Hs.27693	peptidylprolyl isomerase (cyclophilin)-like 1
	107130 RC_AA620582	AB033106	Hs.12913	KIAA1280 protein
		-		

	407450 00 44004000	4.4407040		and the state of t
	107156 RC_AA621239	AA137043	Hs.9663	programmed cell death 6-interacting protein
	107174 RC_AA621714	BE122762	Hs.25338	ESTs
	130621 RC_AA621718	AW513087	Hs.16803	LUC7 (S. cerevislae)-like
5	107190 RC_D19673	AA836401	Hs.5103	ESTs .
)	132626 RC_D25755_s	AW504732	Hs.21275	hypothetical protein FU11011
	107217 RC_D51095	AL080235	Hs.35861	DKFZP586E1621 protein
	131610 RC_D60272_i	AA357879	Hs.29423	scavenger receptor with C-type lectin
	129604 T08879	AF088886	Hs.11590	cathepsin F
10	107295 T34527	AA186629	Hs.80120	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminythransferase 1
10	(GalNAc-T1)	DE033 453	11- 20004	home the flood country by CO 1000
	107299 T40327_s	BE277457	Hs.30661	hypothetical protein MGC4606
	107315 T62771_s	AA316241	Hs.90691	nucleophosmin/nucleoplasmin 3
	107316 T63174_s '	T63174	Hs.193700	Homo sapiens mRNA; cDNA DKFZp586I0324 (from clone DKFZp586I0324)
1.5	107328 T83444	AW959891	Hs.76591	KIAA0887 protein
15	107334 T93641	T93597	Hs.187429	ESTs
	134715 U48263	U48263	Hs.89040	prepronociceptin
	128636 U49065	U49065	Hs.102865	interleukin 1 receptor-like 2
	129938 U79300	AW003668	Hs.135587	Human clone 23629 mRNA sequence
20	107375 U88573	BE011845	Hs.251064	high-mobility group (nonhistone chromosomal) protein 14
20	130074 U93867	AL038596	Hs.250745	polymerase (RNA) III (DNA directed) (62kD)
	107387 W01094	D86983	Hs.118893	Melanoma associated gene
	132036 W01568	AL157433	Hs.37706	hypothetical protein DKFZp434E2220
	107426 W26853	W26853	Hs.291003	hypothetical protein MGC4707
25	113857 W27179	AW243158	Hs.5297	DKFZP564A2416 protein
23	135388 W27965	W27965	Hs.99865	epimorphin
	130419 W36280_s	AF037448	Hs.155489	NS1-associated protein 1
	107469 W47063	W47063	Hs.94668	ESTs
	132616 W79060	BE262677	Hs.283558	hypothetical protein PRO1855
20	107506 W88550	AB028981	Hs.8021	KIAA1058 protein
30	132358 X60486	NM_003542	Hs.46423	H4 histone family, member G
	107522 X78931_s	X78931	Hs.99971	zinc finger protein 272
,	125827 Z14077_s	NM_003403	Hs.97496	YY1 transcription factor
	107582 RC_AA002147	AA002147	Hs.59952	EST hyperficial protein CL 142602
35	107609 RC_AA004711	R75654	Hs.164797	hypothetical protein FLJ13693
22	107661 RC_AA010383	AA010383	Hs.60389	ESTS
	107714 RC_AA015761	AA015761	Hs.60642	ESTs
	107775 RC_AA018772	AW008846	Hs.60857	ESTs gb:ze66c11.s1 Soares retina N2b4HR Homo saplens cDNA clone IMAGE:363956 3', mRNA
	107832 RC_AA021473_i	AAU21473		go:zeooci r.s i Soares leuna ivzo-ink nomo sapiens conva cione liviage:200500 3, nikiva
40	sequence.	A14/720572	Un 47504	notes at the control of the control
40	107859 RC_AA024835	AW732573	Hs.47584	potassium voltage-gated channel, delayed-rectifier, subfamily S, member 3
	124337 RC_AA025858	N23541	Hs.281561	Homo saplens cDNA: FLJ23582 fis, clone LNG13759 ESTs, Weakly similar to T16370 hypothetical protein F45E12.5 - Caenorhabditis elegans
	107914 RC_AA027229	AA027229	Hs.61329	ESTS, Wearly Similar to 1 10370 hypothetical protein F43E12.5 - Caenoniaudits elegans
	[C.elegans]	4.6000.400	Un CAEEE	ECT-
45	107935 RC_AA029428	AA029428	Hs.61555	ESTS
43	116262 RC_AA035143	A1936442	Hs.59838	hypothetical protein FLJ10808
	131461 RC_AA035237	AA992841	Hs.27263	KIAA1458 protein
	108007 RC_AA039347	AA039347	Hs.61916	EST
	108029 RC_AA040740	AA040740	Hs.62007	ESTs. SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily b,
50	108040 RC_AA041551	AL121031	Hs.159971	SyryShr retated, matrix associated, actin dependent regulator of Chromatin, Subiamily 0,
20	member 1	A A 0.E.D.O.A.A	Hs.116602	Hama contrary dans IMACE: 4154009 mDNA portiol ads
	108084 RC_AA045513	AA058944		Homo sapiens, clone IMAGE:4154008, mRNA, partial cds
	108088 RC_AA045745	AA045745	Hs.62886 Hs.63176	ESTs ESTs
	108168 RC_AA055348	Al453137		
55	130719 RC_AA056582_9		Hs.14235	hypothetical protein FLJ20008; KIAA1839 protein ESTs, Moderately similar to A46010 X-linked retinopathy protein [H.saplans]
55	108189 RC_AA056697	AW376061	Hs.63335	
	108190 RC_AA056746		Hs.63338	EST Hamping aDNA+EL 121522 for along COLOGOAD
	108203 RC_AA057678		Hs.289005	Homo sapiens cDNA: FLJ21532 fis, clone COL06049
	108216 RC_AA058681	AA524743	Hs.44883	ESTs
60	108217 RC_AA058686	AA058686	Hs.62588	ESTs
00	108245 RC_AA062840	BE410285	Hs.89545	proteasome (prosome, macropain) subunit, beta type, 4 gb:zm50f03.s1 Stratagene fibroblast (937212) Homo sapiens cDNA clone IMAGE:529085 3',
	108277 RC_AA064859	AA064859		guizinoulos i Sualagene nationast (93/212) nonto sapiens corre done inviociozados a,
	mRNA	AAGCEGCO		shumidanid of Ciminana passage (027200) Hama capiana albita dana 21 milkita capiana
	108280 RC_AA065069	AA065069		gb:zm12e11.s1 Stratagene pancreas (937208) Homo sapiens cDNA clone 3', mRNA sequence gb:zm57e03.r1 Stratagene neuroepithelium (937231) Homo sapiens cDNA clone 5' similar to
65	108309 RC_AA069923	AA069818	No 278270	
05	133739 RC_AA070799_s		Hs.278270	unactive progesterone receptor, 23 kD
	108340 RC_AA070815	AA069820	Hs. 180909	peroxiredoxin 1 gb:zm87a01.s1 Stratagene ovarian cancer (937219) Homo sapiens cDNA done IMAGE:544872
	108403 RC_AA075374	AA075374		guzimiran (3) onamgene ovalian canten (35/213) i ionio sapiena corea cone invidez 34/0/2
	3', mRNA sequence.	A A070300		about 01/00 of Chalanges species consect (027740) Home contens child class IMACT-E4E240
70	108427 RC_AA076382	AA076382		gb:zm91g08.s1 Stratagene ovarian cancer (937219) Homo sapiens cDNA clone IMAGE:545342
70	3', mRNA sequence.	T00407	He 40/404	Home contact of NA: EL MOSCO Se dans ADVANOSTY
	108435 RC_AA078787	T82427	Hs.194101	Homo sapiens cDNA: FLJ20869 fis, done ADKA02377
	108439 RC_AA078986	AA078986		gb:zm92h01.s1 Stratagene ovarian cancer (937219) Homo sapiens cDNA clone IMAGE:545425
	3', mRNA sequence.	A & 0.70000	No 2462	adachema e avidaca minurit VIIIe
75	108465 RC_AA079393	AA079393	Hs.3462	cytochrome c oxidase subunit VIIc gb:zm97f08.s1 Stratagene colon HT29 (937221) Homo sapiens cDNA done 3*, mRNA
13	108469 RC_AA079487	AA079487		Spreng ino 2 to standard mini Li Ca (2017 Truth adhena cristy cand 2 tultur
	sequence			

	108500 RC_AA083207 108501 RC_AA083256		Hs.68270	EST gb:zn08g12.s1 Stratagene hNT neuron (937233) Homo saplens cDNA clone 3' similar to
_	gb:M33308 108533 RC_AA084415	AA084415		gb:zn06g09.s1 Stratagene hNT neuron (937233) Homo saptens cDNA clone IMAGE:546688 3',
5	mRNA 108562 RC_AA085274	AA100796		gb:zm26c06.s1 Stratagene pancreas (937208) Homo sapiens cDNA clone 3' similar to
	gb:X15341 108589 RC_AA088678		Hs.68846	ESTs
10	130890 RC_AA100925 134585 RC_AA101255		Hs.76698 Hs.278573	stress-associated endoplasmic reticulum protein 1; ribosome associated membrane protein 4 H-2K binding factor-2
	130385 RC_AA126474 108749 RC_AA127017	AW067800	Hs.155223 Hs.71052	stanniocalcin 2 ESTs
	108807 RC_AA129968 108808 RC_AA130240	AI652236	Hs.49376 Hs.62738	hypothetical protein FLJ20844 ESTs
15	108833 RC_AA131866 107290 RC_AA132039	AF188527	Hs.61661 Hs.323780	ESTs, Weakly similar to AF174605 1 F-box protein Fbx25 [H.sapiens] ESTs
	108846 RC_AA132983 108857 RC_AA133250	AL117452	Hs.44155 Hs.62180	DKFZP586G1517 protein anillin (Drosophila Scraps homolog), actin binding protein
	131474 RC_AA133583		Hs.2726	high-mobility group (nonhistone chromosomal) protein isoform I-C
20	108894 RC_AA135941		Hs.5105	hypothetical protein FLJ10569
	108941 RC_AA148650 IMAGE:567202 3',			gb:zo09e06.s1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens cDNA clone
	108968 RC_AA151110 108996 RC_AA155754	AW995610	Hs.188680 Hs.332436	ESTS EST
25	109001 RC_AA156125		Hs.72116	hypothetical protein FLJ20992 similar to hedgehog-interacting protein
	131183 RC_AA156289 109019 RC_AA156997		Hs.285107 Hs.72150	· hypothetical protein FLJ13397 ESTs
	109013 RC_AA150397	AA157291	Hs.21479	ubinuclein 1
	109023 RC_AA157293	AA157293	Hs.72168	ESTs
30	109068 RC_AA164293		Hs.72545	ESTs
	109072 RC_AA164676 129021 RC_AA167375		Hs.22394 Hs.173081	hypothetical protein FLJ10893 KIAA0530 protein
	130346 RC_AA167550		Hs.188757	Homo sapiens, clone MGC:5564, mRNA, complete cds
	109146 RC_AA176589		Hs.142078	EST
35	109172 RC_AA180448		Hs.144300	EST
	131080 RC_AA187144		Hs.2271	endothelin 1
	129208 RC_AA189170 109222 RC_AA192757		Hs.109441 Hs.333512	MSTP033 protein similar to rat myomegalin
	109300 RC_AA205650		Hs.170142	ESTs
40	109481 RC_AA233342 109485 RC_AA233472		Hs.289069 Hs.28465	hypothetical protein FLJ21016 Homo sapiens cDNA: FLJ21869 fis, clone HEP02442
	109516 RC_AA234110		Hs.71913	ESTs
	109537 RC_D80981	AI858695	Hs.34898	ESTs
45	109556 RC_F01660 109577 RC_F02206	A1925294 F02206	Hs.87385 Hs.296639	ESTs Homo sapiens potassium channel subunit (HERG-3) mRNA, complete cds
73	109578 RC_F02208	F02208	Hs.27214	ESTs
	109595 RC_F02544	AA078629	Hs.27301	ESTs
	109625 RC_F03918	H29490	Hs.22697	ESTs
50	131983 RC_F04258_s 109648 RC_F04600	AF119665 H17800	Hs.184011 Hs.7154	pyrophosphatase (inorganic) ESTs
50	109671 RC_F08998	R59210	Hs.26634	ESTs
	109699 RC_F09605	H18013	Hs.167483	ESTs
	109820 RC_F11115	AW016809	Hs.323795	ESTs
55	109933 RC_H06371 110014 RC_H10995	R52417	Hs.20945	Homo sapiens clone 24993 mRNA sequence Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 35907
55	110039 RC_H11938	AL109666 H11938	Hs.7242 Hs.21907	histone acetyltransferase
	110099 RC_H16568	R44557	Hs.23748	ESTs
	110107 RC_H16772	AW151660	Hs.31444	ESTs
60	110155 RC_H18951 110197 RC_H20859	A1559526 AW090386	Hs.93522	Homo sapiens mRNA for KIAA1647 protein, partial cds
00	110197 RC_H20039 110223 RC_H23747	H19836	Hs.112278 Hs.31697	arrestin, beta 1 ESTs
	110306 RC_H38087	H38087	Hs.105509	CTL2 gene
	110335 RC_H40331	H65490	Hs.18845	ESTs
65	110342 RC_H40567	H40961	Hs.33008	ESTs
65	110395 RC_H46966 110511 RC_H56640_j	AA025116 H56640	Hs.33333 Hs.221460	ESTS ESTS
	110523 RC_H57154	AI040384	Hs.19102	ESTs, Weakly similar to organic anion transporter 1 [H.sapiens]
	110715 RC_H96712	H96712	Hs.269029	ESTs
70	110754 RC_N20814	AW302200	Hs.6336	KIAA0672 gene product
70	130132 RC_N25249	U55936 NM 016569	Hs.184376 Hs.267182	synaptosomal-associated protein, 23kD TBX3-iso protein
	131135 RC_N27100 134263 RC_N39616	NM_016569 AW973443	Hs.8086	RNA (guanine-7-) methyttransferase
	110938 RC_N48982	N48982	Hs.38034	Homo sapiens cDNA FLJ12924 fis, clone NT2RP2004709
a c	110983 RC_N51957	NM_015367	Hs.10267	MIL1 protein
75	115062 RC_N52271	AA253314	Hs.154103	LIM protein (similar to rat protein kinase C-binding enigma)
	111081 RC_N59435	Al146349	Hs.271614	CGI-112 protein

	:444400 BQ 1/04400	*********	11. 40074	1 and the second of the second
	111128 RC_N64139	AW505364	Hs.19074	LATS (large tumor suppressor, Drosophila) homolog 2
	135244 RC_N66981	AI834273 AW139408	Hs.9711 Hs.152940	novel protein ESTs
	111216 RC_N68640 437562 RC_N69352	AB001636	Hs.5683	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 15
5	131002 RC_N95226	AL050295	Hs.22039	KIAA0758 protein
•	111399 RC_R00138	AW270776	Hs.18857	ESTs
	111514 RC_R07998	R07998		gb:yf16g11.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA done IMAGE:127076 3'
	similar to			
10	130182 RC_R08929	BE267033	Hs.192853	ubiquitin-conjugating enzyme E2G 2 (homologous to yeast UBC7)
10	111574 RC_R10307	Al024145	Hs.188526	ESTs
	111804 RC_R33354	AA482478	Hs.181785	ESTs
	111831 RC_R36083	R36095	Hs.268695	ESTs
	129675 RC_R37938_f 111904 RC_R39330	NM_015556 Z41572	Hs.172180	KIAA0440 protein gb:HSCZYB122 normalized infant brain cDNA Homo saptens cDNA clone c-zyb12, mRNA
15	sequence	2413/2		903 1002 1 0122 1011 lanced than t dails out of 10110 sapiets color daile o'2)012, linder
	133868 RC_R40816_s	AB012193	Hs.183874	CUIIIN 4A
	112033 RC_R43162_s	R49031	Hs.22627	ESTs
	130987 RC_R45698	BE613269	Hs.21893	hypothetical protein DKFZp761N0624
20	112300 RC_R54554	H24334	Hs.26125	ESTS
20	112513 RC_R68425 112514 RC_R68568	R68425 R68568	Hs.13809 Hs.183373	hypothetical protein FLJ10648 src homology 3 domain-containing protein HIP-55
	112522 RC_R68763	R68857	Hs.265499	ESTs
	112540 RC_R70467	R69751	1 10.200 100	gb:yi40a10.s1 Soares placenta Nb2HP Homo sapiens cDNA clone 3', mRNA sequence
	130346 RC_R73565	H05769	Hs.188757	Homo sapiens, done MGC:5564, mRNA, complete cds
25	129534 RC_R73640	AK002126	Hs.11260	hypothetical protein FLJ11264
	112597 RC_R78376	R78376	Hs.29733	EST
	112732 RC_R92453	R92453	Hs.34590	ESTs
	131458 RC_T03865	BE297567	Hs.27047	hypothetical protein FLJ20392
30	112888 RC_T03872 131863 RC_T10072	AW195317 Al656378	Hs.107716 Hs.33461	hypothetical protein FLJ22344 ESTs
50	112911 RC_T10080	AW732747	Hs.13493	like mouse brain protein E46
	132215 RC_T10132	AL035703	Hs.4236	KIAA0478 gene product
	112931 RC_T15343	T02966	Hs.167428	ESTs
25	112984 RC_T23457	T16971	Hs.289014	ESTs, Weakly similar to A43932 mucin 2 precursor, Intestinal [H.sapiens]
35	112998 RC_T23555	H11257	Hs.22968 Hs.7232	Homo sapiens clone IMAGE:451939, mRNA sequence acetyl-Coenzyma A carboxylase alpha
	133376 RC_T23670 113026 RC_T23948	BE618768 AA376654	Hs.183684	eukaryotic translation initiation factor 4 gamma, 2
	113070 RC_T33464	AB032977	Hs.6298	KIAA1151 protein
	128970 RC_T34413	Al375672	Hs.165028	ESTs
40	113074 RC_T34611	AK001335	Hs.31137	protein tyrosine phosphatase, receptor type, E
	113095 RC_T40920	AA828380	Hs.126733	ESTs
	113179 RC_T55182 113337 RC_T77453	BE622021 T77453	Hs.152571 Hs.302234	ESTs, Highly similar to IGF-II mRNA-binding protein 2 [H.saplens] ESTs
	113421 RC_T84039	A1769400	Hs.189729	ESTs
45	113454 RC_T86458	AJ022166	Hs.16188	ESTs
	113481 RC_T87693	T87693	Hs.204327	EST
	131441 RC_T89350_s	AA302862	Hs.90063	neurocalcin delta
	113557 RC_T90945	H66470	Hs.16004	ESTs
50	113559 RC_T90987 113589 RC_T91863	T79763 AI078554	Hs.14514 Hs.15682	ESTs ESTs
50	113591 RC_T91881	T91881	Hs.200597	KIAA0563 gene product
	113619 RC_T93783_s	R08665	Hs.17244	hypothetical protein FLJ 13605
	113683 RC_T96687	AB035335	Hs.144519	T-cell leukemia/lymphoma 6
<i></i>	113692 RC_T96944	AL360143	Hs.17936	DKFZP434H132 protein
55	113702 RC_T97307	T97307		gb:ye53h05.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:121497 3',
	mRNA 113717 RC_T97764	T99513	Hs.187447	ESTs
	113824 RC_W48817	AI631964	Hs.34447	ESTs
	113840 RC_W58343	R72137	Hs.7949	DKFZP586B2420 protein
60	113844 RC_W59949	Al369275	Hs.243010	Homo sapiens cDNA FLJ14445 fis, clone HEMBB1001294, highly similar to GTP-BINDING
	PROTEIN TC10	4.4040444	11- 400000	and Array and Array Array Array 1
	113902 RC_W74644 113904 RC_W74761	AA340111 AF125044	Hs.100009 Hs.19196	acyl-Coenzyme A oxidase 1, palmitoyl ubiquitin-conjugating enzyme HBUCE1
•	113905 RC_W74802	R81733	Hs.33106	ESTs
65	113931 RC_W81205	BE255499	Hs.3496	hypothetical protein MGC15749
	113932 RC_W81237	AA256444	Hs.126485	hypothetical protein FLJ12604; KIAA1692 protein
	131965 RC_W90146_f	W79283	Hs.35962	ESTs
	114035 RC_W92798	W92798	Hs.269181	ESTS ab-DC6 BT0662 260400 044 A02 BT0662 Home conjunction at NA mPNA conjunction
70	114106 RC_Z38412 133593 RC_Z38709	AW602528 Al416988	Hs.238272	gb:RC5-BT0562-260100-011-A02 BT0562 Homo sapiens cDNA, mRNA sequence inositol 1,4,5-triphosphate receptor, type 2
, 0	114161 RC_Z38904	BE548222	Hs.299883	hypothetical protein FLJ23399
	424949 RC_Z39103	AF052212	Hs.153934	core-binding factor, runt domain, alpha subunit 2; translocated to, 2
	129059 RC_Z39930_f	AW069534	Hs.279583	CGI-81 protein
75	128937 RC_Z39939	AA251380	Hs.10726	ESTS, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
75	WARNING	AJ479813	Hs.278411	NCK-associated protein 1
	130983 RC_Z40012_i	(UT) 30 (3	i loci Utili	Houseward Morest I

[C.elegans] 114304 RC_Z40820 At934204 Hs.16129 ESTs 114364 RC_Z41680 AL117427 Hs.172778 Homo sapiens mRNA; cDNA DKFZp566P013 (from clone DKF 5 132900 RC_AA005112 AA777749 Hs.5978 LIM domatin only 7 129034 RC_AA005432 AA481157 Hs.108110 DKFZP547E2110 protein	2 - Caenorhabditis elegans
114364 RC_Z41680 AL117427 Hs.172778 Homo sapiens mRNA; cDNA DKFZp566P013 (from clone DKF 132900 RC_AA005112 AA777749 Hs.5978 LIM domain only 7 129034 RC_AA005432 AA481157 Hs.108110 DKFZP547E2110 protein	
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	•
131881 RC_AA010163 AW361018 Hs.3383 upstream regulatory element binding protein 1	
452461 RC_A0026356 N78223 Hs.108106 transcription factor	
114465 RC_AA026901 BE621056 Hs.131731 hypothetical protein FLJ11099 10 131376 RC_AA036867 AK001644 Hs.26156 hypothetical protein FLJ10782	
101567 RC_AA044644 M33552 Hs.56729 lysosomal	
431555 RC_AA046426 Al815470 Hs.260024 Cdc42 effector protein 3	
132944 RC_AA054515 T96641 Hs.6127 Homo sapiens cDNA: FLJ23020 fis, clone LNG00943	
114618 RC_AA084162 AW979261 Hs.291993 ESTs	
15 130274 RC_AA085749 AA128376 Hs.153884 ATP binding protein associated with cell differentiation	
110330 RC_AA098874 Al288666 Hs.16521 DKFZP434I116 protein 114648 RC_AA101056 AA101056 gbzzn25b03.s1 Stratagene neuroepithelium NT2RAMI 937234	Homo saniens dDNA done
IMAGE:548429 3'	Tions supuls contrations
114658 RC AA102746 AA102383 Hs.249190 tumor necrosis factor receptor superfamily, member 10a	
20 132456 RC_AA114250_s AB011084 Hs.48924 KIAA0512 gene product; ALEX2	
131319 RC_AA126561_s NM_003155 Hs.25590 stanniocalcin 1	Illeria de la composição
132225 RC_AA128980_i AA128980 gb:zo09a11.s1 Stratagene neuroepithelium NT2RAMI 937234 IMAGE:567164 3'	Homo sapiens cuna cione
132669 RC_AA129757 W38586 Hs.293981 guanine nucleotide binding protein (G protein), gamma 3, linke	d
25 114709 RC_AA129921 AA397651 Hs.301959 profine synthetase co-transcribed (bacterial homolog)	•
131973 RC_AA133331 AB018284 Hs.158688 KIAA0741 gene product	
114750 RC_AA135958 AA887211 Hs.129467 ESTs	
115714 RC_AA136524_s T19228 Hs.172572 hypothetical protein FLJ20093	
114763 RC_AA147044 AA810755 Hs.88977 hypothetical protein dJ511E16.2 30 114767 RC_AA148885 Al859865 Hs.154443 minichromosome maintenance deficient (S. cerevisiae) 4	
114774 RC_AA150043 AV656017 Hs.184325 CGI-76 protein	
129388 RC_AA151621 AA662477 Hs.110964 hypothetical protein FLJ23471	
129183 RC_AA155743 BE561824 Hs.273369 uncharacterized hematopoletic stem/progenitor cells protein M	DS027
128869 RC_AA156335 AA768242 Hs.80618 hypothetical protein 35 130207 RC_AA156336 AF044209 Hs.144904 nuclear receptor co-repressor 1	
35 130207 RC_AA156336 AF044209 Hs.144904 nuclear receptor co-repressor 1 114798 RC_AA159181 AA159181 Hs.54900 serologically defined colon cancer antigen 1	
114800 RC_AA159825 Z19448 Hs.131887 ESTs, Weakly similar to T24396 hypothetical protein T03F6.2	- Caenorhabditis elegans
[C.elegans]	
114828 RC_AA234185 AA252937 Hs.283522 Homo saplens mRNA; cDNA DKFZp434J1912 (from clone DK	FZp434J1912)
40 114846 RC_AA234929 BE018682 Hs.166196 ATPase, Class I, type 8B, member 1	
114848 RC_AA234935 BE614347 Hs.169615 hypothetical protein FLJ20989 114902 RC_AA236359 AW275480 Hs.39504 hypothetical protein MGC4308	
132271 RC_AA236466 AB030034 Hs.115175 sterile-alpha motif and leucine zipper containing kinase AZK	
114907 RC_AA236535 N29390 Hs.13804 hypothetical protein dJ462O23.2	
45 135159 RC_AA236935_s U43374 Hs.95631 Human normal keratinocyte mRNA	
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	126884 RC	_AA279667_s		Hs.297939 Hs.286236	cathepsin B
	115322 RC		U49436 L08895	Hs.78995	KIAA1856 protein MADS box transcription enhancer factor 2, polypeptide C (myocyte enhancer factor 2C)
	133626 RC		AW836130	Hs.75277	hypothetical protein FLJ13910
5	115372 RC		AW014385	Hs.88678	ESTs, Weakly similar to Unknown [H.saplens]
•		_AA283127_s		Hs.57698	Empirically selected from AFFX single probeset
			F05422	Hs.168352	nucleoporin-like protein 1
	129192 RC		AA286914	Hs.183299	ESTs
	452598 RC		AI831594	Hs.68647	ESTs, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE CONTAMINATION
10	WARNING	_			
	132131 RC	_AA293495	AF069291	Hs.40539	chromosome 8 open reading frame 1
	115536 RC	_AA347193	AK001468	Hs.62180	anillin (Drosophila Scraps homolog), actin binding protein
	132411 RC	_AA398474_s	AA059412	Hs.47986	hypothetical protein MGC10940
	115575 RC	_AA398512	AA393254	Hs.43619	ESTs
15	115601 RC	_AA400277	AA148984	Hs.48849	ESTS, Weakly similar to ALU4_HUMAN ALU SUBFAMILY SB2 SEQUENCE CONTAMINATION
	Warning				
				Hs.199160	myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog)
	125819 RC			Hs.251871	CTP synthase
20	115683 RC			Hs.54650	junctional adhesion molecule 2
20		_		Hs.1390	proteasome (prosome, macropain) subunit, beta type, 2
	132952 RC	_	A1658580	Hs.61426	Homo saplens mesenchymal stem cell protein DSC96 mRNA, partial cds
	115819 RC		AA486620	Hs.41135	endomucin-2
	132525 RC			Hs.50727	N-acetylgtucosaminidase, alpha- (Santilippo disease IIIB)
25			AB033035	Hs.51965	KIAA1209 protein
23				Hs.45032	ESTS
				Hs.179520	hypothetical protein MGC10702
		_		Hs.42911	ESTS
	115974 RC			Hs.238944	hypothetical protein FLJ10631 ECTA Wooldy similar to T09500 prohable transcription factor CA150 (Magnison)
30	115985 RC			Hs.268115 Hs.1098	ESTs, Weakly similar to T08599 probable transcription factor CA150 [H.sapiens]
50				Hs.64313	DKFZp434J1813 protein ESTs, Weakly similar to AF257182 1 G-protein-coupled receptor 48 [H.sapiens]
	116095 RC		AA043429	Hs.62618	ESTs
		_AA460454_s		Hs.172788	ALEX3 protein
				Hs.172788	ALEX3 protein
35				Hs.326740	hypothetical protein MGC10947
	134585 RC		D14041	Hs.278573	H-2K binding factor-2
	134790 RC			Hs.287850	Integral membrane protein 1
	116265 RC			Hs.55189	hypothetical protein
		_AA485084_s		Hs.4947	hypothetical protein FLJ22584
40		_AA485431_8		Hs.182874	guanine nucleotide binding protein (G protein) alpha 12
	303150 RC	_AA489057	AA887146	Hs.8217	stromal antigen 2
	129945 RC	_AA489638	BE514376	Hs.165998	PAI-1 mRNA-binding protein
	116331 RC		N41300	Hs.71616	Homo sapiens mRNA; cDNA DKFZp586N1720 (from clone DKFZp586N1720)
46 .	116333 RC			Hs.203963	hypothetical protein FLJ10339 .
45	132994 RC			Hs.279905	done HQ0310 PRO0310p1
				Hs.85951	exportin, tRNA (nuclear export receptor for tRNAs)
	116391 RC			Hs.75113	general transcription factor IIIA
		_AA599574_j		Hs.65370	lipase, endothelial
50				Hs.110713	DEK oncogene (DNA binding)
20	116417 RC			Hs.12484	Human clone 23826 mRNA sequence
	116429 RC			Hs.279923	putative nucleotide binding protein, estradio-induced
	116439 RC 116459 RC			Hs.43913 Hs.302738	PIBF1 gene product Homo sapiens cDNA: FLJ21425 fis, done COL04162
	427505 RC			Hs.178761	26S proteasome-associated pad1 homolog
55	132699 RC			Hs.55200	ESTs
55	116541 RC			Hs.249212	polymerase (RNA) III (DNA directed) (155kD)
	132557 RC			Hs.5122	ESTs
	112259 RC			Hs.333402	hypothetical protein MGC12760
	116571 RC		D45652		gb:HUMGS02848 Human adult lung 3' directed Mool cDNA Homo sapiens cDNA 3', mRNA
60	sequence.				
	129815 RC	_D60208_f	BE565817	Hs.26498	hypothetical protein FLJ21657
	421919 RC		AJ224901	Hs.109526	zinc finger protein 198
	116643 RC	_F03010	Al367044	Hs.153638	myeloid/lymphoid or mixed-lineage leukemia 2
	116661 RC	_F04247	R61504		gb:yh16a03.s1 Soares infant brain 1NIB Homo sapiens cDNA clone 3' similar to contains Alu
65	repetitive				
	116715 RC	_F10966	AL117440	Hs.170263	tumor protein p53-binding protein, 1
	116729 RC			Hs.115823	ribonuclease P, 40kD subunit
	318709 RC		R52576	Hs.285280	Hamo saptens cDNA: FLJ22096 fis, done HEP16953
70	134760 RC				erythropoletin receptor
70	116773 RC			Hs.169149	karyopherin alpha 1 (importin alpha 5)
	106425 RC			Hs.247423	adducin 2 (beta)
	116780 RC	_FIZZ300		Hs.30098	ESTs
	131978 RC	_IT40433_\$		Hs.36232	KIAA0186 gene product
75	116819 RC			Hs.93698	EST VIA ANGON
13	111428 RC 133175 RC			Hs.174174	KIAA0601 protein
	1331/3 RU	_ N: 30! _8	AW955632	Hs.66666	ESTs, Wealthy similar to S19560 proline-rich protein MP4 - mouse [M.muscutus]

	116844 RC_H64938_s 116845 RC_H64973 116892 RC_H69535	H64938 AA649530 AI573283	Hs.337434 Hs.38458	ESTs, Weakly similar to A46010 X-linked retinopathy protein [H.sapiens] gb:ns44f05.s1 NCL_CGAP_Ahr1 Homo saptens cDNA clone, mRNA sequence ESTs
_	116925 RC_H73110	H73110	Hs.260603	ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.saptens]
5	116981 RC_H81783 131768 RC_H86259	N29218 AC005757	Hs.40290 Hs.31809	ESTs . hypothetical protein
	117031 RC_H88353 contains L1	H88353		gb:yw21a02.s1 Morton Fetal Cochlea Homo sapiens cDNA done IMAGE:252842 3' similar to
10	117034 RC_H88639 132542 RC_H88675	U72209 AL137751	Hs.180324 Hs.263671	YY1-associated factor 2 Homo saplens mRNA; cDNA DKFZp434l0812 (from clone DKFZp434l0812); partial cds
	134403 RC_H93708_s 117280 RC_N22107	AA334551 M18217	Hs.82767 Hs.172129	sperm spediic antigen 2 Homo saglens cDNA: FLJ21409 fis, clone COL03924
	117344 RC_N24046	R19085	Hs.210706	Homo sapiens cDNA FLJ13182 fis, clone NT2RP3004070
15	117422 RC_N27028	Al355562 N30205	Hs.43880	ESTs, Weakly similar to A46010 X-linked retinopathy protein [H,sapiens] ESTs, Weakly similar to 138022 hypothetical protein [H.sapiens]
13	117475 RC_N30205 117487 RC_N30621	N30621	Hs.93740 Hs.44203	ESTs Treatily suiting to 100022 hypothetical protein (n.sapiens)
	130207 RC_N33258	AF044209	Hs.144904	nuclear receptor co-repressor 1 EST
	117549 RC_N33390 117683 RC_N40180	N33390 N40180	Hs.44483	gb;yy44d02.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone
20	IMAGE:276387 3' similar to	o N45198	Hs.47248	ESTs, Highly similar to similar to Cdc14B1 phosphatase [H.sapiens]
	117710 RC_N45198 104514 RC_N45979_s	AF164622	Hs.182982	golgin-67
	117791 RC_N48325	N48325	Hs.93956	ĒST EST-
25	117822 RC_N48913 129647 RC_N49394	AA706282 AB018259	Hs.93963 Hs.118140	ESTs KIAA0716 gene product
	117895 RC_N50656 [H.sapiens]	AW450348	Hs.93996	ESTs, Highly similar to SORL_HUMAN SORTILIN-RELATED RECEPTOR PRECURSOR
	131557 RC_N50721	AA317439	Hs.28707	signal sequence receptor, gamma (translocon-associated protein gamma)
30	133057 RC_N53143 118103 RC_N55326	AA465131 AA401733	Hs.64001 Hs.184134	Hamo sapiens clone 25218 mRNA sequence ESTs
50	118111 RC_N55493	N55493	113,104134	gb:yv50c02.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:246146 3',
	mRNA 118129 RC_N57493	N57493		gb:yy54c08.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone
25	IMAGE:277358 3', mRNA		11 040400	
35	118278 RC_N62955 118329 RC_N63520	N62955 N63520	Hs.316433	Homo sapiens cDNA FLJ11375 fis, clone HEMBA1000411, weakly similar to ANKYRIN gb;yy62f01.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone IMAGE:278137
	3', mRNA	•	U- 47400	•
	118336 RC_N63604 132457 RC_N64166	BE327311 AB017365	Hs.47166 Hs.173859	HT021 frizzled (Drosophila) homolog 7
40	118363 RC_N64168	Al183838	Hs.48938	hypothetical protein FLJ21802
	118364 RC_N64191 118475 RC_N66845	N46114 N66845	Hs.29169	hypothetical protein FLJ22623 gb:za46c11.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:295604 3'
	similar to 118491 RC_N67135	AV647908	Hs.90424	Hamo sapiens cDNA: FLJ23285 fis, clone HEP09071
45	118500 RC_N67295	W32889	Hs.154329	ESTs
	101663 RC_N68399 118584 RC_N68963	NM_003528 AW136928	Hs.2178	H2B histone family, member Q gb:UI-H-BI1-adp-d-08-0-UI.s1 NCI_CGAP_Sub3 Homo sapiens cDNA done 3', mRNA
	sequence		Un 440004	
50	421983 RC_N69331 118661 RC_N70777	Al252640 AL137554	Hs.110364 Hs.49927	peptidytprolyl isomerase C (cyclophilin C) protein kinase NYD-SP15
	118684 RC_N71364_s	N71313	Hs.163986	Homo sapiens cDNA: FLJ22765 fts, clone KAIA1180
	118689 RC_N71545_s 118690 RC_N71571	AW390601 N71571	Hs.184544 Hs.269142	Homo saplens, clone IMAGE:3355383, mRNA, partial ods ESTs
55	118766 RC_N74456	N74456	Hs.50499	EST
23	118793 RC_N75594 118817 RC_N79035	N75594 AI668658	Hs.285921 Hs.50797	ESTs, Moderately similar to T47135 hypothetical protein DKFZp761L0812.1 [H.sapiens] ESTs
	118844 RC_N80279	AL035364	Hs.50891	hypothetical protein
	118919 RC_N91797 129558 RC_N92454	AW452696 AW580922	Hs.130760 Hs.180446	myosin phosphatase, target subunit 2 karyopherin (Importin) beta 1
60	132692 RC_N94581	AW191962	Hs.249239	collagen, type VIII, alpha 2
	118996 RC_N94746 119021 RC_N98238	N94746 N98238	Hs.274248 Hs.55185	hypothetical protein FLJ20758 ESTs
	119039 RC_R02384	Al160570	Hs.252097	pregnancy specific beta-1-glycoprotein 6 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
65	119063 RC_R16833 WARNING	R16833	Hs.53106	ESTS, Moderately similar to ALO I_HOWAN ALO SOBFAMILT 3 SECULIACE CONTAMINATION
	118523 RC_R41828_s	Y07759 T02865	Hs.170157 Hs.328321	myosin VA (heavy polypeptide 12, myoxin) EST
	119111 RC_R43203 133970 RC_R46395	AA214228	Hs.127751	hypothetical protein ·
70	119146 RC_R58863 120296 RC_R78248	R58863 AW995911	Hs.91815 Hs.299883	ESTs hypothetical protein FLJ23399
70	119239 RC_T11483	T11483	. 13.200000	gb:CHR90049 Chromosome 9 exon Homo saplens cDNA clone 111-1 5 and 3, mRNA
	sequence. 119281 RC_T16896	Al692322	Hs.65373	ESTs, Wealthy similar to T02345 hypothetical protein KIAA0324 [H.sapiens]
75	119298 RC_T23820	NM_001241	Hs.155478	cyclin T2
75	126502 RC_T30222 135073 RC_W15275_s	T10077 W55956	Hs.13453 Hs.94030	hypothetical protein FLJ14753 Homo sapiens mRNA; cDNA DKFZp586E1624 (from clone DKFZp586E1624)

	440750 70 1170404	1400404		E II B I I I I I I I I I I I I I I I I I
	119558 RC_W38194	W38194	11- 000004	Empirically selected from AFFX single probeset
	132736 RC_W42414_s sapiens mad protein	AW081883	Hs.288261	Homo sapiens cDNA: FLJ23037 fis, done LNG02036, highly similar to HSU68019 Homo
	132173 RC_W46577_s	X89426	Hs.41716	endothelial cell-specific molecula 1
5	134873 RC_W49632_s	AA884471	Hs.90449	Human clone 23908 mRNA sequence
•	119650 RC_W57613	R82342	Hs.79856	ESTs, Weakly similar to \$65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
	119654 RC_W57759	W57759		gb:zd20g11.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:341252 3'
	similar to			
10	119683 RC_W61118	W65379	Hs.57835	ESTs
10	119694 RC_W65344	AA041350	Hs.57847	ESTs, Moderately similar to ICE4_HUMAN CASPASE4 PRECURSOR [H.sapiens]
	119718 RC_W69216	W69216	Hs.92848	ESTS Home position mBNA: mDNA DVET/nE98D0022 /from close DVET/nE98D0022\
	133010 RC_W69379 119938 RC_W86728	Al287518 AW014862	Hs.62669 Hs.58885	Homo sapiens mRNA; cDNA DKFZp586D0923 (from clone DKFZp586D0923) ESTs
	120128 RC_Z38499	BE379320	Hs.91448	MKP-1 like protein tyrosine phosphatase
15	120130 RC_Z38630	AA045767	Hs.5300	bladder cancer associated protein
	120148 RC_Z39494	F02806	Hs.65765	ESTs
	120155 RC_Z39623	Z39623	Hs.65783	ESTs
	131486 RC_Z40071_s	F06972	Hs.27372	BMX non-receptor tyrosine kinase
20	120183 RC_Z40174	AW082866	Hs.65882	ESTs
20	120184 RC_Z40182	Z40182	Hs.65885	EST .
	120211 RC_Z40904	Z40904	Hs.66012	EST
	120245 RC_AA166965	AW959615	Hs.111045	ESTs
	120247 RC_AA167500 120254 RC_AA169599_8	AA167500	Hs.103939 Hs.111054	EST ESTs
25	120259 RC_AA171724	AW014786	Hs.192742	hypothetical protein FLJ12785
	120260 RC_AA171739	AK000061	Hs.101590	hypothetical protein
	120275 RC_AA177105	AA177105	Hs.78457	solute carrier family 25 (mitochondrial carrier; ornithine transporter) member 15
	120284 RC_AA182626	AA179656		gb:zp54e11.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone 3' similar
	to contains			
30	114056 RC_AA186324	AA188175	Hs.82506	KIAA1254 protein
	129507 RC_AA192099	AJ236885	Hs.112180	zinc finger protein 148 (pHZ-52)
	120302 RC_AA192173	AA837098	Hs.269933	ESTs
	120303 RC_AA192415 120305 RC_AA192553	Al216292 AW295096	Hs.96184	ESTS
35	120305 RC_AA192555 120319 RC_AA194851	T57776	Hs.101337 Hs.191094	uncoupling protein 3 (mitochondrial, proton carrier) ESTs
33	133389 RC_AA195520_s		Hs.72639	ESTs
	120326 RC_AA196300	AA196300	Hs.21145	hypothetical protein RG083M05.2
	134272 RC_AA196517	X76040	Hs.278614	protease, serine, 15
	133145 RC_AA196549	H94227	Hs.6592	Homo sapiens, clone IMAGE:2961368, mRNA, partial cds
40	120327 RC_AA196721	AK000292	Hs.278732	hypothetical protein FLJ20285
	106686 RC_AA196729_i		Hs.334825	Homo sapiens cDNA FLJ14752 fis, clone NT2RP3003071
	120328 RC_AA196979	AA923278	Hs.290905	ESTs, Weakly similar to protease [H.sapiens]
	120340 RC_AA206828 similar to	AA206828		gb:zq80b08.s1 Stratagene hNT neuron (937233) Homo sapiens cDNA clone IMAGE:647895 3"
45	134292 RC_AA207123	Al906291	Hs.81234	immunoglobulin superfamily, member 3
	131522 RC_AA214539_i		Hs.239489	TIA1 cytotoxic granule-associated RNA-binding protein
	129051 RC_AA226914_s	AA227068	Hs.108301	nuclear receptor subfamily 2, group C, member 1
	120375 RC_AA227260	AF028706	Hs.111227	Zic tamily member 3 (odd-paired Drosophila homolog, heterotaxy 1)
50	120376 RC_AA227469	AA227469		gb:zr18a07.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone
50	IMAGE:663732 3', mRNA		Un 4444CO	soletum (solenestulin denendent ambie kinoco (CoM kinoco) II dolta
	120390 RC_AA233122 303876 RC_AA233334_s	AA837093	Hs.111460 Hs.66521	calclum/calmodulin-dependent protein kinase (CeM kinase) II delta Machado-Joseph disease (spinocerebellar ataxia 3, olivopontocerebellar ataxia 3, autosomal
	dominant, ataxin 3)	004020	NS.0002 I	maxiado-boseph disease (spilitocerenella) ataxia 3, ottroportuocerenella ataxia 3, admissimal
	132038 RC_AA233347	AI825842	Hs.3776	zinc finger protein 216
55	104463 RC_AA233519	T85825	Hs.246885	hypothetical protein FLi20783
	125750 RC_AA233714	AA018515	Hs.264482	Homo sapiens mRNA; cDNA DKFZp761A0411 (from clone DKFZp761A0411)
	120396 RC_AA233796	AA134006	Hs.79306	eukaryotic translation initiation factor 4E
	120409 RC_AA235050_f	AA235050		gb:zs38e04.s1 Soares_NhHMPu_S1 Homo saplens cDNA clone IMAGE:687486 3' similar to
60	gb:L07077	414407450	H- 404000	handhallani amiala El 140000
60	120414 RC_AA235704	AW137156	Hs.181202	hypothetical protein FLJ10038 spinal cord-derived growth factor-B
	120420 RC_AA236031 120422 RC_AA236352	AI128114 AL133097	Hs.112885 Hs.301717	hypothetical protein DKFZp434N1928
	132221 RC_AA236390_s		Hs.42419	ESTs
	120423 RC_AA236453	AA236453	Hs.18978	Homo sapiens cDNA: FLJ22822 fis, clone KAIA3968
65	120435 RC_AA243370	AA243370	Hs.96450	EST
	120453 RC_AA250947	AA250947	Hs.170263	tumor protein p53-binding protein, 1
	120455 RC_AA251083	AA251720	Hs.104347	ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C WARNING ENTRY !!! [H.sapiens]
	120456 RC_AA251113	AA488750	Hs.88414	BTB and CNC homology 1, basic leucine zipper transcription factor 2
70	120473 RC_AA251973	AA251973	Hs.269988	ESTS
70	128922 RC_AA252023 120477 RC_AA252414	A1244901 AA252414	Hs.9589 Hs.43141	ubiquiim 1 DKFZP727C091 protein
	120477 RC_AA252650	AF006689	Hs.110299	mitogen-activated protein kinase kinase 7
	120488 RC_AA255523	AW952916	Hs.63510	KIAAD141 gene product
	120510 RC_AA258128	AI796395	Hs.111377	ESTs
75	120527 RC_AA262105	AA262105	Hs.4094	Homo sapiens cDNA FLJ14208 fis, clone NT2RP3003264
	120528 RC_AA262107	Al923511	Hs.104413	ESTs

	400000 BO 4400000	A140.4000	11- 404445	FOT-
	120529 RC_AA262235	AI434823	Hs.104415	ESTs
	120541 RC_AA278298	W07318	Hs.240	M-phase phosphoprotein 1
	131445 RC_AA278529_i	_	Hs.172052	serine/ihreonine kinase 18
5	120544 RC_AA278721	BE548277	Hs.103104	ESTS
5	120562 RC_AA280036	BE244580	Hs.302267	hypothetical protein FLJ10330 ESTs, Weakly similar to B34323 GTP-binding protein Rab2 [H.saplens]
	120569 RC_AA280648	AA807544	Hs.24970	
	120571 RC_AA280738	AB037744	Hs.34892	KIAA1323 protein
	120572 RC_AA280794	H39599	Hs.294008	ESTs
10	129434 RC_AA280837	AW967495	Hs.186644	ESTs gbzp39e03.s1 Stratagene muscle 937209 Homo saplens cDNA clone 3' similar to contains Alu
10	130529 RC_AA280886	AA178953		gozposedo.s i Sualagene musice sor zos monto sapiens corea cone o suntan lo contano ano
	repetitive	AM/070000	Un 220044	hundhalled ambin DVE7a76254544; VIAA4946 ambin
	120575 RC_AA280934	AW978022	Hs.238911 Hs.54037	hypothetical protein DKFZp762E1511; KIAA1816 protein
	132635 RC_AA281535	AB020686		ectorucleotide pyrophosphatase/phosphodiesterase 4 (putative function) general transcription factor IIH, polypeptide 2 (44kD subunit)
15	120591 RC_AA281797_s		Hs.191356 Hs.193522	general danscription raction first, polypepide 2 (44kD subdility) ESTs
13	120593 RC_AA282047 430275 RC_AA283002	Z11773	Hs.237786	zinc finger protein 187
	117729 RC_AA283709	AA306166	Hs.7145	calpain 7
	120609 RC_AA283902	AW978721	Hs.266076	ESTs, Weakly similar to A46010 X-linked retinopathy protein [H.sapiens]
	132754 RC_AA284108	AI752244	Hs.75309	eukaryotic translation elongation factor 2
20	130315 RC_AA284109	Al241084	Hs.154353	nonselective sodium potassium/proton exchanger
20	132614 RC_AA284371	AA284371	Hs.118064	similar to rat nuclear ubiquitous casein kinase 2
	447503 RC_AA284744_f		Hs.336898	Homo sapiens, Similar to RIKEN cDNA 1810038N03 gene, clone MGC:9890, mRNA, complete
	cds	701110450	113.555555	There are the second of the se
	135376 RC_AA284784	BE617856	Hs.99756	mitochondrial ribosome recycling factor
25	120621 RC_AA284840	AW961294	Hs.143818	hypothetical protein FLJ23459
	107868 RC_AA286844	AA286844	Hs.61260	hypothetical protein FLJ13164
	129868 RC_AA287032		Hs.13012	ESTs
	120644 RC_AA287038	Al869129	Hs.96616	ESTs
	120660 RC_AA287546	AA286785	Hs.99677	ESTs
30	135370 RC_AA287553_s	BE622187	Hs.99670	ESTs, Weakly similar to I38022 hypothetical protein [H.sapiens]
	120661 RC_AA287556		Hs.263412	ESTs, Weakly similar to ALUB_HUMAN !!!! ALU CLASS B WARNING ENTRY !!! [H.sapiens]
	129116 RC_AA287564	AB019494	Hs.225767	IDN3 protein
	131567 RC_AA291015_s		Hs.28853	CDC7 (cell division cycle 7, S. cerevisiae, homolog)-like 1
	120699 RC_AA291716	A1683243	Hs.97258	ESTs, Moderately similar to S29539 ribosomal protein L13a, cytosolic [H.sapiens]
35	100690 RC_AA291749_s		Hs.1657	estrogen receptor 1
	120726 RC_AA293656	AA293655	Hs.97293	ESTs
	120737 RC_AA302430	AL049176	Hs.82223	chordin-like
	120745 RC_AA302809	AA302809		gb:EST10426 Adipose tissue, white I Homo saplens cDNA 3' end, mRNA sequence.
40	135192 RC_AA302820_s		Hs.321709	purinergic receptor P2X, ligand-gated ion channel, 4
40	120750 RC_AA310499	AI191410	Hs.96693	ESTs, Moderately similar to 2109260A B cell growth factor (H.sapiens)
	120761 RC_AA321890	AA321890	Hs.1265	branched chain keto acid dehydrogenase E1, beta polypeptide (maple syrup urine disease)
	120768 RC_AA340589	AA340589	Hs.104560 Hs.96769	EST ESTs
	120769 RC_AA340622 135232 RC_AA342457_i	AI769467	Hs.96800	ESTS, Moderately similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE
45	CONTAMINATION	ALU30012	F13.30000	LOTS, WANDEREDY SHIMME TO ALLOY TO MAIN ALLO DO DE ANNIET DE CERCETOLE
73	133439 RC_AA342828_s	723091	Hs.73734	glycoprotein V (platelet)
	120793 RC_AA342864	AA342864	Hs.96812	ESTs
	120796 RC_AA342973	Al247356	Hs.96820	ESTs
	120809 RC_AA346495	AA346495		gb:EST52657 Fetal heart II Homo sapiens cDNA 3' end similar to EST containing O family
50	repeat, mRNA sequence.			
•	132459 RC_AA347573	AL120071	Hs.48998	fibronectin leucine rich transmembrane protein 2
	120825 RC_AA347614	AI280215	Hs.96885	ESTs
	120827 RC_AA347717	AA382525	Hs.132967	Human EST clone 122887 mariner transposon Hsmar1 sequence
	120839 RC_AA348913	AA348913		gb:EST55442 Infant adrenal gland II Homo sapiens cDNA 3' end similar to EST containing Alu
55	repeat, mRNA sequence.			
	120850 RC_AA349647		Hs.96927	Homo sapiens cDNA FLJ12573 fis, clone NT2RM4000979
	120852 RC_AA349773		Hs.191564	ESTs
	128852 RC_AA350541_s		Hs.106601	ESTs
C 0	135240 RC_AA357159_i		Hs.96986	EST
60	120870 RC_AA357172_j	AA357172	Hs.292581	ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
	WARNING	1107000	11- 400044	
	134637 RC_AA369856_s		Hs.180941	vacuolar protein sorting 41 (yeast homolog)
	120894 RC_AA370132		Hs.97063	ESTS
65	131854 RC_AA370472_s		Hs.173202	L-kappa-B-interacting Ras-like protein 1 ESTs, Moderately similar to AF174605 1 F-box protein Fbx25 [H.sapiens]
05	120897 RC_AA370867		Hs.97079 Hs.97104	ESTs, inductately stituted to AF (74000 11-000 protest) to AED (1504) and 1505
	120915 RC_AA377296 120935 RC_AA383902			ESTS. Weakly similar to ALU1 HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
	WARNING	CUPUPUU	Hs.97177	CO10, FROMING WALCOT, HORIZITATE COOR ARREST O CERCENCE CONTINUMENTOR
	120936 RC_AA385934	AA385034	Hs.97184	EST, Highly similar to (defline not available 7499603) [C.elegans]
70	120937 RC_AA386255		Hs.97186	EST
7.5	120938 RC_AA386260		Hs.104632	EST
	129722 RC_AA386266	R20855	Hs.5422	glycoprotein M6B
	120950 RC_AA398014	AA398014	Hs.104684	EST
		AI219896	Hs.97592	ESTs
75	120988 RC_AA398235	AA398235	Hs.97631	ESTs

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121008 RC_AA398348 AA398348
                                             Hs.301720
                                                          Human DNA sequence from clone RP11-251J8 on chromosome 13 Contains ESTs, STSs,
        GSSs and a CpG
         121029 RC_AA398482
                                AA398482
                                             Hs.97641
         121032 RC_AA398504
                                AA393037
                                             Hs.161798
                                                          ESTs
 5
         121033 RC_AA398505
                                AA398505
                                             Hs.97360
                                                          ESTs
         121034 RC_AA398507
                                AL389951
                                             Hs.271623
                                                           nucleoporin 50kD
         121035 RC_AA398523
                                AA398523
                                             Hs.210579
                                                          ESTs
         121058 RC_AA398625
                                AA398625
                                             Hs.97391
                                                          ESTs
         121060 RC_AA398632
                                AA398632
                                             Hs.97395
                                                          ESTs
10
         121061 RC_AA398633
                                AA393288
                                             Hs.97396
                                                          ESTs
         121091 RC_AA398894
                                AA398894
                                             Hs.97657
                                                          ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE
        CONTAMINATION
         121092 RC_AA398895
                                AA398895
                                             Hs.97658
                                                          gb:zt62h10.r1 Soares_testis_NHT Homo sapiens cDNA clone 5', mRNA sequence
        121094 RC_AA398900
                                AA402505
15
         121096 RC_AA398904
                                AA398904
                                             Hs.332690
                                                          ESTs
         121115 RC_AA399122
                                AA398187
                                             Hs.104682
                                                          ESTs, Wealdy similar to mitochondrial citrate transport protein [H.sapiens]
         121121 RC_AA399371
                                AA399371
                                             Hs.189095
                                                          similar to SALL1 (sal (Drosophlla)-like
        121122 RC_AA399373
                                AJ126713
                                             Hs.192233
                                                          ESTs, Highly similar to T00337 hypothetical protein KIAA0568 [H.sapiens]
        121125 RC_AA399441
121151 RC_AA399636
                                                          KIAA1201 protein
                                AL042981
                                             Hs.251278
20
                                AA399636
                                             Hs.143629
                                                          ESTs
        121153 RC_AA399640
                                AA399640
                                             Hs.97694
                                                          ESTs
         121163 RC_AA399680
                                AI676062
                                             Hs.111902
                                                          ESTs
         121176 RC_AA400080
                                AL121523
                                             Hs.97774
                                                          ESTs
        121192 RC_AA400262
121223 RC_AA400725
                                AA400262
                                             Hs.190093
                                                          ESTs
25
                                Al002110
                                             Hs.97169
                                                          ESTs, Weakly similar to dJ667H12.2.1 [H.sapiens]
        121227 RC_AA400748
                                                           Homo sapiens mRNA; cDNA DKFZp434D024 (from clone DKFZp434D024)
                                AA400748
                                             Hs.97823
                                                          ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C WARNING ENTRY !!! [H.sapiens]
        121231 RC_AA400780
                                AA814948
                                             Hs.96343
        121278 RC_AA401631
                                             Hs.98518
                                                          Homo sapiens cDNA FLJ11490 fis, clone HEMBA1001918
                               AA037121
        121279 RC_AA401688
121282 RC_AA401695
                                AA292873
                                             Hs.177996
30
                                AA401695
                                             Hs.97334
                                                          ESTs
        121299 RC_AA402227
                                AA402227
                                             Hs.22826
                                                          tropomodulin 3 (ubiquitous)
        121301 RC_AA402329
                                NM_006202
                                             Hs.89901
                                                           phosphodiesterase 4A, cAMP-specific (dunce (Drosophila)-homolog phosphodiesterase E2)
         121302 RC_AA402398
                                AA402587
                                             Hs.325520
                                                           LAT1-3TM protein
        121304 RC_AA402449
                                AA293863
                                             Hs.97316
35
        121305 RC_AA402468
                                AA402468
                                             Hs.291557
                                                          ESTs
         134721 RC_AA403268_s AK000112
                                             Hs.89306
                                                          hypothetical protein FLJ20105
        121323 RC_AA403314
                               AA291411
                                             Hs.97247
                                                          ESTs
        121324 RC_AA404229
                                AA404229
                                             Hs.97842
                                                          EST
         129047 RC_AA404260
                                AI768623
                                             Hs.108264
                                                          ESTs
40
        131074 RC_AA404271
                                U16125
                                             Hs.181581
                                                          glutamate receptor, ionotropic, kalnate 1
        121344 RC_AA405026
121348 RC_AA405182
121350 RC_AA405237
                                AA405026
                                             Hs.193754
                                                          ESTs
                                AA405182
                                             Hs.97973
                                AA405237
                                                          gb:zt06e10.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:712362 3' similar to
        contains Alu
45
         121400 RC_AA406061
                                AA406061
                                             Hs.98001
        121402 RC_AA406063
121403 RC_AA406070
                                AA406063
                                             Hs.98003
                                                          ESTs
                                AA406070
                                             Hs.98004
                                                          EST
        121408 RC_AA406137
                                AA406137
                                             Hs.98019
                                                          EST
        121431 RC_AA406335
                                AA035279
                                             Hs.176731
                                                          ESTs
50
         132936 RC_AA411801
                                AL120659
                                             Hs.6111
                                                          aryl-hydrocarbon receptor nuclear translocator 2
         121471 RC_AA411804
                                AA411804
                                             Hs.261575
                                                          ESTs
        121474 RC_AA411833
                                AA402335
                                             Hs.188760
                                                          ESTs, Highly similar to Trad [H.sapiens]
        121526 RC_AA412219
121530 RC_AA412259
121558 RC_AA412497
                                AW665325
                                             Hs.98120
                                                          ESTs
                                AA778658
                                             Hs.98122
                                                          ESTs
55
                                AA412497
                                                          gb:zt95g12.s1 Soares_testis_NHT Homo sapiens cDNA clone IMAGE:730150 3' similar to
        contains L1.13 L1
        121559 RC_AA412498
                                Al192044
                                             Hs.104778
                                                          ESTs
        121584 RC_AA416586
                                AI024471
                                             Hs.98232
                                                          ESTs
        121609 RC_AA416867
                                             Hs.98185
                                AA416867
                                                          EST
60
        121612 RC_AA416874
                                AA416874
                                             Hs.98168
                                                          ESTs
        121737 RC_AA421133
                               AA421133
                                             Hs.104671
                                                          erythrocyte transmembrane protein
        121740 RC_AA421138
                                AA421138
                                             Hs.98334
        129194 RC_AA422079
                                             Hs.109276
                                AA150797
                                                          latexin protein
        121784 RC_AA423837
                                T90789
                                             Hs.94308
                                                          RAB35, member RAS oncogene family
65
        121802 RC_AA424328
                                             Hs.188898
                                Al251870
                                                          ESTs
        121803 RC_AA424339
                                AI338371
                                             Hs.157173
                                                          ESTs
         135286 RC_AA424469_s AW023482
                                             Hs.97849
                                                          ESTs
                                AA424313
        121806 RC_AA424502
                                             Hs.98402
                                                          ESTs
        129517 RC_AA425004
                                AW972853
                                             Hs.112237
                                                          ESTs
70
        121845 RC_AA425734
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                                             Hs.165066
                                                          ESTs, Moderately similar to ALU2_HUMAN ALU SUBFAMILY SB SEQUENCE
        CONTAMINATION
                                AA425887
                                             Hs 98502
                                                          hypothetical protein FLJ14303
        121853 RC_AA425887
        121891 RC_AA426456
                                AA426456
                                             Hs.98469
        121895 RC_AA427396
                                AA427396
                                                          gb:zw33a02.s1 Soares ovary turnor NbHOT Homo sapiens cDNA clone IMAGE:771050 3*
75
        similar to contains
        121899 RC_AA427555
                               R55341
                                             Hs.50421
                                                          KIAA0203 gene product
```

```
121917 RC_AA428218
                                AA406397
                                             Hs.98038
                                                          ESTs
        121918 RC_AA428242
                                BE274689
                                             Hs.184175
                                                          chromosome 2 open reading frame 3
        121919 RC AA428281
                                AA428281
                                             Hs.98560
                                                          EST
        121941 RC_AA428865
                                AA428865
                                             Hs.98563
                                                          ESTs
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        121942 RC_AA428994
                                AW452701
                                             Hs.293237
                                                          ESTs
        121970 RC_AA429666
                                AA429666
                                             Hs.98617
                                                          EST
        121993 RC_AA430181
                                AW297880
                                             Hs.98661
                                                          ESTs
        134660 RC_AA430184_s U73524
                                             Hs 87465
                                                          ATP/GTP-binding protein
                                                          CD3D antigen, delta polypeptide (TiT3 complex)
        126753 RC_AA431288_s AA306478
                                             Hs.95327
10
                                                          ESTs, Moderately similar to T42650 hypothetical protein DKFZp434D0215.1 [H.sapiens]
        122022 RC_AA431293
                                AA431293
                                             Hs.98716
        122050 RC_AA431478
                                A1453076
                                             Hs.166109
                                                          ELAV (embryonic lethal, abnormal vision, Drosophila)-like 2
        122051 RC_AA431492
122055 RC_AA431732
                                AA431492
                                             Hs.98742
                                                          EST
                                AA431732
                                             Hs.98747
                                                          EST
         122105 RC_AA432278
                                             Hs.98699
                                AW241685
                                                          ESTs
15
        122125 RC_AA434411
                                AK000492
                                             Hs.98806
                                                          hypothetical protein
        135235 RC_AA435512_I AW298244
                                             Hs.293507
                                             Hs.79946
                                                          cytochrome P450, subfamily XIX (aromatization of androgens)
        122162 · RC_AA435698
                                AA628233
        129406 RC_AA435711
                                AB018255
                                             Hs.111138
                                                          KIAA0712 gene product
        318801 RC_AA435815_s U40763
                                             Hs.77965
                                                          peptidyl-prolyt isomerase G (cyclophilin G)
20
        122186 RC_AA435842
                                AA398811
                                             Hs.104673
                                                          ESTs
        122235 RC_AA436475
                                AA436475
                                             Hs.112227
                                                          membrane-associated nucleic acid binding protein
                                                          dual specificity phosphatase 10
        129131 RC_AA436489
                                AB026436
                                             Hs.177534
        134664 RC_AA442060
                                AA256106
                                             Hs.87507
                                                          ESTs
         122310 RC_AA442079
                                                          ESTs, Wealthy similar to S65824 reverse transcriptase homolog [H.sapiens]
                                AW192803
                                             Hs.98974
                                                          ESTs, Weakly similar to LB4D_HUMAN NADP-DEPENDENT LEUKOTRIENE B4 12-
25
                                             Hs.98365
        122334 RC_AA443151
                                BE465894
        122382 RC_AA446133
                                AA446440
                                             Hs.98643
        122425 RC_AA447145
122431 RC_AA447398
                                AB007859
                                             Hs.100955
                                                          KIAA0399 protein
                                AA447398
                                             Hs.99104
                                                          ESTs
                                                          hypothetical protein DKFZp434F1819
         122450 RC_AA447643
                                             Hs.112095
                                AA447643
30
                                                          dynein, axonemal, heavy polypeptide 9
        302653 RC_AA447742_s AJ404468
                                             Hs.284259
        122477 RC_AA448226
                                AA448226
                                             Hs.324123
                                                          ESTs
        122500 RC_AA448825
                                AA448825
                                             Hs.99190
                                                          ESTs
        122522 RC_AA449444
                                AA299607
                                             Hs.98969
                                                          ESTs
        122536 RC_AA450087
                                AF060877
                                             Hs.99236
                                                          regulator of G-protein signalling 20
35
        122538 RC_AA450211
                                AA450211
                                             Hs.99239
                                                          ESTs, Wealthy similar to A43932 mucin 2 precursor, Intestinal [H.sapiens]
         122540 RC_AA450244
                                AA476741
                                             Hs.98279
         122560 RC_AA452123
                                AW392342
                                             Hs.283077
                                                          centrosomal P4.1-associated protein; uncharacterized bone marrow protein BM032
                                AJ224901
        421919 RC_AA452155
                                             Hs.109526
                                                          zinc finger protein 198
                                                          gb:zx29c03.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:787876 3',
        122562 RC_AA452156
                                AA452156
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        mRNA.
        122585 RC_AA453036
                                A1681654
                                             Hs.170737
                                                          hypothetical protein FLJ23251
        122608 RC_AA453526
                                AA453525
                                             Hs.143077
        122635 RC_AA454085
                                AA454085
                                                          gb:zx33a08.s1 Soares_total_fetus_Nb2HF8_9w Homo sapiens cDNA clone IMAGE:788246 3'
        similar to
45
        122636 RC_AA454103
                                AW651706
                                             Hs.99519
                                                          hypothetical protein FLJ14007
        122653 RC_AA454642
                                AW009166
                                             Hs.99376
                                                          EST<sub>8</sub>
        122660 RC_AA454935
                                A1816827
                                             Hs.180069
                                                          nuclear respiratory factor 1
         122703 RC_AA456323
                                AA456323
                                             Hs.269369
                                                          ESTs
        122724 RC_AA457395
                                AA457395
                                             Hs.99457
                                                          ESTs
50
        122749 RC_AA458850
                                             Hs.293372
                                AA458850
                                                          ESTs, Weakly similar to B34087 hypothetical protein [H.sapiens]
        122772 RC_AA459662
                                             Hs.99489
                                AW117452
                                                          ESTs
                                                          3-hydroxyisobutyryi-Coenzyme A hydrolase hypothetical protein FLJ13409; KIAA1711 protein
        131098 RC_AA459668
                                U66669
                                             Hs.236642
        129045 RC_AA459679_s Al082883
                                             Hs.30732
                                                          hypothetical protein FLJ10160 similar to insulin related protein 2
        122777 RC_AA459702
                                AK001022
                                             Hs.214397
55
        135362 RC_AA460017_f AA978128
                                             Hs.99513
                                                          ESTs, Weakly similar to T17454 diaphanous-related formin - mouse [M.musculus]
                                AW366286
        122798 RC_AA460324
                                             Hs.145696
                                                          splicing factor (CC1.3)
        122837 RC_AA461509
                                             Hs.293565
                                                          ESTs. Wealty similar to putative p150 [H.sapiens]
                                AA461509
        122860 RC_AA464414_i AA464414
                                                          gb:zx78g01.s1 Soares ovary turnor NbHOT Homo sapiens cDNA clone IMAGE:809904 3*,
        mRNA sequence. ·
60
        122861 RC_AA464428
                                AA335721
                                             Hs.119394
        122910 RC_AA470084
                                AA470084
                                             Hs.98358
                                                          ESTs
        132899 RC_AA476606_s AA476606
                                             Hs.59666
                                                          SMAD in the antisense orientation
        122967 RC_AA478521
                                AA806187
                                             Hs.289101
                                                          glucose regulated protein, 58kD
                                                          hypothetical protein MGC2752
        129560 RC_AA478523
                                AA317841
                                             Hs.7845
65
        123009 RC_AA479949
                                AA535244
                                             Hs.78305
                                                          RAB2, member RAS oncogene family
        128917 RC_AA481252
                                Al365215
                                             Hs.206097
                                                          oncogene TC21
        123081 RC_AA485351
                                Al815486
                                             Hs.243901
                                                          Homo sapiens cDNA FLJ20738 fis, clone HEP08257
        123133 RC_AA487264
                                             Hs.154974
                                                          Homo sapiens mRNA; cDNA DKFZp667N064 (from clone DKFZp667N064)
                                AA487264
         123184 RC_AA489072
                                BE247767
                                             Hs 18166
                                                          KIAA0870 protein
70
         129671 RC_AA489630
                                             Hs.119004
                                                          KIAA0665 gene product
                                NM_014700
                                                          ESTs, Wealty similar to MAPB_HUMAN MICROTUBULE-ASSOCIATED PROTEIN 1B
         123233 RC_AA490225
                                AW974175
                                             Hs.188751
        [H.sapiens]
         123234 RC_AA490227
                                                          down-regulator of transcription 1, TBP-binding (negative cofactor 2)
                                NM_001938
                                             Hs.16697
        123236 RC_AA490255
                                AW968504
                                             Hs.123073
                                                          CDC2-related protein kinase 7
75
                                                          ESTs
         123255 RC_AA490890
                                AA830335
                                             Hs 105273
         129503 RC_AA490916_s AW768399
                                             Hs.112157
                                                          ESTs
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	404040 50 44400505	. ===		
	131043 RC_AA490925	AF084535	Hs.22464	epilepsy, progressive myodonus type 2, Lafora disease (laforin)
	123259 RC_AA490955 [H.sapiens]	AI744152	Hs.283374	ESTs, Weakly similar to CA15_HUMAN COLLAGEN ALPHA 1(V) CHAIN PRECURSOR
	123284 RC_AA495812	AA488988	Hs.293796	ESTs
5	123286 RC_AA495824	AA495824	Hs.188822	ESTs, Weakly similar to A46010 X-linked retinopathy protein (H.sapiens)
_	123315 RC_AA496369	AA496369		gb:zx/37d10.s1 Soares ovary tumor NbHOT Homo saplens cDNA clone IMAGE:755827 3' similar
	to contains			6
	129179 RC_AA504125_s	AW969025	Hs.109154	ESTs
10	131612 RC_AA521473	AU076668	Hs.334884	SEC10 (S. cerevistae)-like 1
10	123421 RC_AA598440	AA598440	Hs.291154	EST, Weakly similar to 138022 hypothetical protein [H.sapiens]
	123449 RC_AA598899_I	AL049325 AL044675	Hs.112493	Homo sapiens mRNA; cDNA DKFZp564D036 (from clone DKFZp564D036)
	129021 RC_AA599244 132830 RC_AA599694_s		Hs.173081 Hs.57730	KIAA0530 protein KIAA0133 gene product
	123497 RC_AA600037	AA765256	Hs.135191	ESTs, Weakly similar to unnamed protein product [H.saplens]
15	123604 RC_AA609135	AA609135	Hs.293076	ESTs
	129539 RC_AA609582	T47614	Hs.323022	ESTs, Highly similar to p60 katanin (H.sapiens)
	123712 RC_AA609684	AA609684	Hs.112748	Homo sapiens cDNA: FLJ21543 fis, clone COL06171
	123731 RC_AA609839	AA609839		gb:ae62f01.s1 Stratagene lung cardinoma 937218 Homo sapiens cDNA clone IMAGE:951481 3*
20	similar to	********	U= 00040	CALA bis disc and the gard with anything anticing
20	130725 RC_AA609862 123800 RC_AA620423	T98807 AA620423	Hs.80248 Hs.112862	RNA-binding protein gene with multiple splicing EST
	123841 RC_AA620747	AA620747	Hs.112896	ESTs
	123929 RC_AA621364	AA621364	Hs.112981	ESTs
	123978 RC_C20653	T89832	Hs.170278	ESTs
25	133184 RC_D20085	AA001021	Hs.6685	thyrold hormone receptor interactor 8
	132835 RC_D20749	Z83844	Hs.5790	hypothetical protein dJ37E16.5
	132406 RC_D51285_s	AL133731	Hs.4774	Homo sapiens mRNA; cDNA DKFZp761C1712 (from clone DKFZp761C1712)
	128695 RC_D59972_I	NM_003478	Hs.101299	cullin 5 gb:HSC2JH062 normalized infant brain cDNA Homo saplens cDNA clone c-2jh06 3', mRNA
30	124028 RC_F04112_f sequence.	F04112		guinoczanocz nomalized iliani bian corex nomo sapiens corex done czylod 3, mistex
50	124057 RC_F13604	AA902384	Hs.73853	bone morphogenetic protein 2
	134899 RC_H01662	A1609045	Hs.321775	hypothetical protein DKFZp434D1428
	130973 RC_H05135_i	Al638418	Hs.78580	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 1
25	124106 RC_H12245	H12245	U- 404770	gb:ym17a12.r1 Soares infant brain 1NIB Homo sapiens cDNA clone 3', mRNA sequence
35	124136 RC_H22842 124165 RC_H30894	H22842 H30039	Hs.101770 Hs.107674	EST ESTs
	131229 RC_H43442_s	NM_015340	Hs.2450	leucyl-tRNA synthetase, mitochondrial
	124178 RC_H45996	BE463721	Hs.97101	putative G protein-coupled receptor
	129948 RC_H69281_j	AI537162	Hs.263988	ESTs
40	134374 RC_H69485_f	N22687	Hs.8236	ESTs
	124254 RC_H69899	H69899		gb:yu70c12.s1 Weizmann Olfactory Epithelium Homo sapiens cDNA clone IMAGE:239158 3*
	similar to 129056 RC_H70627_s	Al769958	Hs.108336	ESTs, Weakly similar to ALUE_HUMAN !!!! ALU CLASS E WARNING ENTRY !!! [H.sapiens]
	100919 RC_H73050_s	X54534	Hs.278994	Rhesus blood group, CoEe antigens
45	130724 RC_H73260	AK001507	Hs.306084	Homo sapiens clone FLB6914 PRO1821 mRNA, complete cds
	100716 RC_H77531_s	X89887	Hs.172350	HIR (histone cell cycle regulation defective, S. cerevisiae) homolog A
	124274 RC_H80552	H80552	Hs.102249	EST
	129078 RC_H80737_s	AI351010	Hs.102267	lysosomal
50	124828 RC_H93412 124315 RC_H94892_s	AW952124 NM_005402	Hs.13094 Hs.288757	presenilins associated rhomboid-like protein v-ral simian leukemia viral oncogene homolog A (ras related)
50	100747 RC_H95643_s	X04588	Hs.85844	neurotrophic tyrosine kinase, receptor, type 1
	124324 RC_H96552	H96552	Hs.159472	Homo sapiens cDNA: FLJ22224 fis, done HRC01703
	452933 RC_H97146	AW391423	Hs.288555	Homo sapiens cDNA: FLJ22425 fis, clone HRC08686
<i></i>	132231 RC_H99131_s	AA662910	Hs.42635	hypothetical protein DKFZp434K2435
55	129170 RC_H99462_s	AW250380	Hs.109059	mitochondrial ribosomal protein L12
	133143 RC_H99837_s 132963 RC_N22140	AA094538 AA099693	Hs.272808 Hs.34851	putative transcription regulation nuclear protein; KIAA1689 protein epsilon-tubulin
	135297 RC_N22197	AL118782	Hs.300208	Sec23-interacting protein p125
	134347 RC_N23756_s	AF164142	Hs.82042	solute carrier family 23 (nucleobase transporters), member 1
60	130365 RC_N24134	W56119	Hs.155103	eukaryotic translation initiation factor 1A, Y chromosome
	421642 RC_N24195	AF172066	Hs.106346	refinoic acid repressible protein
	439311 RC_N26739 124383 RC_N27098	BE270668	Hs.151945	mitochondrial ribosomal protein L43
	124387 RC_N27637	N27098 N27637	Hs.102463 Hs.109019	EST ESTs
65	129341 RC_N33090	AJ193519	Hs.226396	hypothetical protein FLJ11126
	129081 RC_N35967	A1364933	Hs.168913	serine/threonine kinase 24 (Ste20, yeast homolog)
	102827 RC_N38959_f	BE244588	Hs.6456	chaperonin containing TCP1, subunit 2 (beta)
	124433 RC_N39069	AA280319	Hs.288840	PRO1575 protein
70	124441 RC_N46441 132338 RC_N48270_f	AW450481 AA353868	Hs.161333 Hs.182982	ESTs
, 0	131403 RC_N48365_s	AI473114	Hs.26455	gogin-67 ESTs
	124466 RC_N51316	R10084	Hs.113319	kinesin heavy chain member 2
	132210 RC_N51499_s	NM_007203	Hs.42322	A kinase (PRKA) anchor protein 2
75	124483 RC_N53976	AI821780	Hs.179864	ESTS
75	124484 RC_N54157 124485 RC_N54300	H66118 AB040933	Hs.285520 Hs.15420	ESTs, Weakly similar to 2109260A B cell growth factor [H.saplens] KIAA1500 protein
	15400 1747140400	, 1000	10.10120	

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124494 RC_N54831
                                N54831
                                              Hs.271381
                                                           ESTs, Weakly similar to 138022 hypothetical protein [H.sapiens]
         129200 RC_N59849
                                N59849
                                              Hs.13565
                                                           Sam68-like phosphotyrostne protein, T-STAR
         124527 RC_N62132
                                N79264
                                              Hs.269104
         124532 RC_N62375
                                              Hs.102731
                                N62375
                                                           EST
 5
         133213 RC_N63138
                                AA903424
                                              Hs.6786
                                                           ESTs
                                D54120
         124539 RC_N63172
                                              Hs.146409
                                                           cell division cycle 42 (GTP-binding protein, 25kD)
         133651 RC_N63772
                                Al301740
                                              Hs.173381
                                                           dihydropyrimidinase-like 2
                                                           ESTs, Weakly similar to 138022 hypothetical protein [H.sapiens]
        129196 RC_N63787
                                RF296313
                                              Hs.265592
                                                            gb:za11c01.s1 Soares fetal liver spieen 1NFLS Homo saplens cDNA clone 3, mRNA sequence
         124575 RC_N6816B
                                N68168
10
                                                           ESTs, Wealdy similar to I38022 hypothetical protein [H.sapiens]
         124576 RC_N68201
                                N68201
                                              Hs.269124
         124577 RC_N68300
                                                           gb:za12g07.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:292380 3',
                                N68300
        mRNA
         124578 RC_N68321
                                N68321
                                              Hs.231500
                                                           EST
        124593 RC_N69575
128501 RC_N75007
                                N69575
                                              Hs.102788
15
                                                           protein containing CXXC domain 2
                                AL133572
                                              Hs.199009
                                                            Homo sapiens dDNA FLJ11918 fis, done HEMBB1000272
         105691 RC_N75542
                                A1680737
                                              Hs.289068
         128473 RC N90066
                                              Hs.100293
                                                           O-linked N-acetylglucosamine (GlcNAc) transferase (UDP-N-acetylglucosamine:polypeptide-N-
                                T78277
         128639 RC_N91246
                                AW582962
                                              Hs.102897
                                                           CGI-47 protein
         124652 RC_N92751
                                                           regulator of nonsense transcripts 2; DKFZP434D222 protein
                                W19407
                                              Hs.3862
20
         133137 RC_N93214_s
                                              Hs.65746
                                AB002316
                                                           KIAA0318 protein
         124671 RC_N99148
                                AK001357
                                              Hs.102951
                                                           Homo saptens cDNA FLJ10495 fis, clone NT2RP2000297, moderately similar to ZINC FINGER
        PROTEIN
        133054 RC_R07876
                                AA464836
                                              Hs.291079
                                                           ESTs. Weakly similar to T27173 hypothetical protein Y54G11A.9 - Caenorhabditts elegans
        [C.elegans]
130410 RC_R10865_f
25
                                J00077
                                              Hs.155421
                                                            alpha-fetoprotein
                                                           gb:ye91c08.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:125102 3'
         124720 RC_R11056
                                R05283
         similar to
         124722 RC_R11488
                                T97733
                                              Hs.185685
                                                            gb.yh31a05.r1 Soares placenta Nb2HP Homo sapiens cDNA clone 5' similar to contains L1
         129961 RC_R22947
                                R23053
30
        repetitive element 128944
                                RC_R23930_s AL137586
                                                                        anaphase-promoting complex subunit 7
                                                           Hs.52763
                                                            hypothetical protein MGC12936
         132965 RC_R26589_f
                                A1248173
                                              Hs.191460
                                                            RAB2, member RAS oncogene family-like
         133740 RC_R37588_s
                                AW162919
                                              Hs.170160
         133074 RC_R37613
                                AL134275
                                              Hs.6434
                                                            hypothetical protein DKFZp761F2014
                                              Hs.141055
                                                            Homo sapiens clone 23758 mRNA sequence
        124757 RC_R38398
                                H11368
35
         124762 RC_R39179_f
                                              Hs.92096
                                                            ESTs, Moderately similar to A46010 X-linked retinopathy protein [H.sapiens]
                                AA553722
         124773 RC_R40923
                                R45154
                                              Hs.106604
                                                           ESTs
                                                            KIAA0328 protein
         135266 RC_R41179
                                R41179
                                              Hs.97393
         131375 RC_R41294_s
                                AW293165
                                              Hs.143134
                                                            ESTs
                                                           early development regulator 2 (homolog of polyhomeotic 2)
         133753 RC_R42307_f
                                NM_004427
                                              Hs.165263
40
         128540 RC_R43189_f
                                AW297929
                                              Hs.328317
                                W38537
                                                            hypothetical protein MGC3040
         124785 RC_R43306
                                              Hs 280740
                                                            hypothetical protein FLJ20736
         124792 RC_R44357
                                R44357
                                              Hs.48712
         124793 RC_R44519
                                R44519
                                                            gbyg24h04.s1 Soares infant brain 1NIB Homo sapiens cDNA done IMAGE:33350 3', mRNA
        sequence.
45
         124799 RC_R45088
                                                           gb:yg38g04.s1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:34896 3', mRNA
                                R45088
        Seculence.
         124812 RC_R47948_i
                                R47948
                                              Hs.188732
         124821 RC_R51524
                                H87832
                                              Hs.7388
                                                            kelch (Drosophila)-like 3
         127274 RC_R54950
                                AW966158
                                              Hs.58582
                                                            Homo sapiens cDNA FLJ12789 fis, clone NT2RP2001947
50
        124835 RC_R55241
                                R55241
                                              Hs.101214
                                                           EST
                                              Hs.101255
        124845 RC_R59585
                                R59585
                                                           ESTs
        124847 RC_R60044
                                              Hs.304177
                                                            Homo sapiens done FLB8503 PRO2286 mRNA, complete cds
                                W07701
                                                            Human DNA sequence from done RP1-304B14 on chromosome 6. Contains a gene for a novel
        440630 RC_R60872
                                BE561430
                                              Hs.239388
        protein and a part of a gene for a novel protein with two is
                                                           oforms. Contains ESTs, STSs, GSSs and a CpG island
55
         124861 RC_R66690
                                R67567
                                              Hs.107110
                                                           ESTs
         130141 RC_R67266_s
                                NM_004455
                                              Hs.150956
                                                           exostoses (multiple)-like 1
         124879 RC_R73588
                                R73588
                                              Hs.101533
                                                           ESTs
         124892 RC_R79403
                                AI970003
                                              Hs.23756
                                                           hypothetical protein similar to swine acylneuraminate lyase
                                              Hs.107815
        124906 RC_R87647
                                H75964
                                                           ESTs
60
                                R93622
         124922 RC_R93622
                                                            eukaryotic translation initiation factor 2, subunit 2 (beta, 38kD)
                                              Hs.12163
        124940 RC_R99599_s
                                AF068846
                                              Hs.103804
                                                           heterogeneous nuclear ribonucleoprotein U (scaffold attachment factor A)
        124941 RC_R99612
                                AJ766661
                                              Hs.27774
                                                           ESTs, Highly similar to AF161349 1 HSPC086 [H.sapiens]
                                                           ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
        124943 RC_T02888
                                AW963279
                                              Hs.123373
        WARNING ENTRY [H.sapiens]
                                T03170
65
         124947 RC_T03170
                                              Hs.100165
                                                           ESTs
                                AW964237
         124954 RC_T10465
                                              Hs.6728
                                                            KIAA1548 protein
        132924 RC_T15418_f
                                U55184
                                              Hs.154145
                                                            hypothetical protein FLJ11585
        133113 RC_T15597_f
132975 RC_T15652_i
                                              Hs,65238
                                BE383768
                                                            95 kDa refinoblastoma protein binding protein; KIAA0661 gene product
                                R43504
                                              Hs.6181
70
         133235 RC_T16898_s
                                AW960782
                                              Hs.6856
                                                            ash2 (absent, small, or homeotic, Drosophila, homolog)-like
        131082 RC_T26644_I
                                              Hs.246218
                                                           Homo sapiens cDNA: FLJ21781 fis, clone HEP00223
                                Al091121
        124980 RC_T40841
                                T40841
                                              Hs.98681
         124984 RC_T47566_i
                                BE313210
                                              Hs.223241
                                                            eukaryotic translation elongation factor 1 delta (guanine nucleotide exchange protein)
                                                            gb:yb77c10.s1 Stratagene ovary (937217) Homo sapiens cDNA clone IMAGE:77202 3' similar
         124991 RC T50116
                                T50116
75
        to similar to SP:VE22_LAMBD P03756 EA22 GENE, mRNA sequence.
                               NM_004477 Hs.203772
         129475 RC_T50145_s
                                                           FSHD region gene 1
```

	425000 DO TENOAS	TE0045	11- 440040	row.
	125000 RC_T58615	T58615	Hs.110640	ESTS
	132932 RC_T59940_f	AW118826	Hs.6093	Homo sapiens cDNA: FLJ22783 fis, done KAIA1993
	129534 RC_T63595 125008 RC_T64891	AK002126 T91251	Hs.11260	hypothetical protein FLJ11264 gb:yd60a10.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone 3', mRNA sequence
5	125009 RC_T64924	T64924	Hs.303046	go.yoooa to.s.1 Soores retai liver spream martis month sapiters contact cone.5, military sequence.
	132940 RC_T64933_r	T79136	Hs.127243	Homo sapiens mRNA for KIAA1724 protein, partial cds
	125017 RC_T68875	T68875	110.127.240	gb:yc30f05.s1 Stratagene liver (937224) Homo saplens cDNA clone IMAGE:82209 3', mRNA
	sequence.			go., woodoo. 1 out agono area (out and a superior out a done and to account out of the and
	125018 RC_T69027	T69027	Hs.57475	sex comb on midleg homolog 1
10	125020 RC_T69924	T69981		gbryc19d031 Stratagene lung (937210) Homo sapiens cDNA clone 5', mRNA sequence
	129891 RC_T70353	AI084813	Hs.13197	ESTs
	134204 RC_T79780_s	AI873257	Hs.7994	hypothetical protein FLJ20551
	125050 RC_179951	AW970209	Hs.111805	ESTs
	125052 RC_T80174_s	T85104	Hs.222779	ESTs, Moderately similar to similar to NEDD-4 [H.saplens]
15	125054 RC_T80622	T80622	Hs.268601	ESTs, Weakly similar to envelope [H.sapiens]
	125063 RC_T85352	T85352		gb:yd82d01.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:114721 3'
			ontains L1 repe	fitive element; mRNA sequence.
	125064 RC_T85373	T85373	antaine MEDO :	gb.yd82f07.s1 Soares fetal liver spieen 1NFLS Homo sapiens cDNA clone IMAGE:114757 3
20		T86284	oniains meks i	epetitive elament;, mRNA sequence. gb:yd77b07.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone 3' similar to contains
20	125066 RC_T86284 Atu repetitive element, mR			go.yor/100/131 Socies retailiver spicest intrus notito sapiens conta come s sinual to contains
	112264 RC_T89579_s	AL045364	Hs.79353	transcription factor Dp-1
	125080 RC_T90360	T90360	Hs.268620	ESTS, Highly similar to ALU6_HUMAN ALU SUBFAMILY SP SEQUENCE CONTAMINATION
	WARNING ENTRY [H.sapi		110.200020	Coro, right) dring to recognish it rico coor rather or observe contrating the
25	125097 RC_T94328_i	AW576389	Hs.335774	EST, Moderately similar to S65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
	125104 RC_T95590	T95590	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	gb:ye40a03.s1 Soares fetal liver spleen 1NFLS Homo saptens cDNA clone 3' similar to
	gb[M10817]IGURRAA Igua		RNA):. mRNA	· ·
	135107 RC_T97257_f	T97257	Hs.337531	ESTs, Moderately similar to 138022 hypothetical protein [H.sapiens]
	129550 RC_T97599_i	AA845462	Hs.124024	deltex (Drosophila) homolog 1
30	125118 RC_T97620	R10606		gb:yl35f11.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:128877 3'
	similar to contains Alu repe			
	125120 RC_T97775	T97775	Hs.100717	EST
	134160 RC_T98152	T98152	Hs.79432	fibrillin 2 (congenital contractural arachnodactyly)
35	125136 RC_W31479	AW962364	Hs.129051	ESTs
33	125144 RC_W37999	AB037742	Hs.24336	KIAA1321 protein Empirically selected from AFFX single probeset
	125150 RC_W38240 104180 RC_W40150	W38240 AA247778	Hs.119155	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 814975
	131987 RC_W45435	AW453069	Hs.3657	activity-dependent neuroprotective protein
	125178 RC_W58202	W93127	Hs.31845	ESTs
40	125180 RC_W58344	W58469	Hs.103120	EST\$
	125182 RC_W58650	AA451755	Hs.263560	ESTs
	130588 RC_W68736	AL030996	Hs.16411	hypothetical protein LOC57187
	125197 RC_W69106	AF086270	Hs.278554	heterochromatin-like protein 1
	133497 RC_W69111	BE617303	Hs.74266	hypothetical protein MGC4251
45	100562 RC_W69385_s	NM_006185	Hs.301512	nuclear mitotic apparatus protein 1
	125639 RC_W69399_s	Z97630	Hs.226117	H1 histone family, member 0
	129232 RC_W69459	R98881	Hs.109655	sex comb on midleg (Drosophila)-like 1
	101495 RC_W72424	W72424	Hs.112405	S100 calcium-binding protein A9 (calgranulin B) ESTs, Weakly similar to TSP2_HUMAN THROMBOSPONDIN 2 PRECURSOR [H.saplens]
50	125209 RC_W72724 125212 RC_W72834	W72724 AA746225	Hs.103174 Hs.103173	ESTs Treaty similar in 1372_notway tricondoctorolly 2 Preconson (insaplets)
50	129132 RC_W73955	BE383436	Hs.108847	hypothetical protein MGC2749
	125223 RC_W74701	Al916269	Hs.109057	ESTs, Weakly similar to ALU5_HUMAN ALLU SUBFAMILY SC SEQUENCE CONTAMINATION
	WARNING ENTRY [H.sapi		. 10.10000	and the first of t
	125225 RC_W76540	W74169	Hs.16492	DKFZP564G2022 protein
55	125228 RC_W79397	AA033982	Hs.110059	ESTs, Weakly similar to 138022 hypothetical protein [H.saplens]
	132393 RC_W85888	AL135094	Hs.47334	hypothetical protein FLJ14495
	125238 RC_W86038	N99713	Hs.109514	ESTs
	125247 RC_W86881	AA694191	Hs.163914	ESTs
CO	129296 RC_W87804	AI051967	Hs.110122	ESTs
60	125263 RC_W88942	AA098878		gb:zn45g10.r1 Stratagene HeLa cell s3 937216 Homo sapiens cDNA clone 5', mRNA sequence
	125266 RC_W90022	W90022	Hs.186809	ESTs, Highly similar to LCT2_HUMAN LEUKOCYTE CELL-DERIVED CHEMOTAXIN 2
	PRECURSOR [H.sapiens]	U91543	Un DECOI	chromodomain helicase DNA binding protein 3
	131321 RC_W92272	NM_007115	Hs.25601 Hs.29352	tumor necrosis factor, alpha-induced protein 6
65	131601 RC_W92764_s 131677 RC_W93040	H05317	Hs.283549	ESTs
05	120837 RC_W93092	BE149656	Hs.306621	Homo sapiens cDNA FLJ11963 fis, clone HEMBB1001051
	125277 RC_W93227	W93227	Hs.103245	EST
	125278 RC_W93523	Al218439	Hs.129998	enhancer of polycomb 1
70	125280 RC_W93659	AI123705	Hs.106932	ESTs
	131856 RC_W94003_s	W93949	Hs.33245	ESTs
	131844 RC_W94401_s	AI419294	Hs.324342	ESTs
	125284 RC_W94688	NM_002666	Hs.103253	perilipin
	313447 RC_W94787_s	AW016321	Hs.82306	destrin (actin depolymenting factor)
75	130799 RC_Z38294_s	AB028945	Hs.12696	cortactin SH3 domain-binding protein
	125289 RC_Z38311	T34530	Hs.4210	Homo sapiens cDNA FLJ13069 fis, clone NT2RP3001752
	128874 RC_Z38465_s	H06245	Hs.106801	ESTs, Weakly similar to PC4259 ferritin associated protein [H.sapiens]

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AW971018
        130966 RC_Z38525_s
                                             Hs.21659
                                                           ESTs
        128875 RC_Z38538_f
                                AB040923
                                             Hs.106808
                                                           kelch (Drosophila)-like 1
        133200 RC_Z38551_s
                                AB037715
                                                           hypothetical protein FLJ10210
                                             Hs.183639
        130158 RC_Z38783_s
                                AB032947
                                             Hs.151301
                                                           Ca2+-dependent activator protein for secretion
 5
                                                           sema domain, immunoglobulin domain (Ig), transmembrane domain (TM) and short cytoplasmic
        125295 RC_Z39113
                                AB022317
                                             Hs.25887
        domain, (semaphorin) 4F
        125298 RC_Z39255_f
                                AW972542
                                             Hs.289008
                                                           Homo saplens cDNA: FLJ21814 ffs, clone HEP01068
        125300 RC_Z39591
                                             Hs.101376
                                Z39591
                                                           EST
        323122 RC_Z39783_s
                                                           Homo sapiens cDNA FLJ12908 fis, clone NT2RP2004399
                                             Hs.264915
                                BE622770
10
        311463 RC_Z39920
                                R55344
                                             Hs.22142
                                                           cytochrome b5 reductase b5R.2
        130882 RC_Z40166_f
                                AA497044
                                             Hs.20887
                                                           hypothetical protein FLJ10392
        128888 RC_Z40388_s
                                A1760853
                                             Hs.241558
                                                           ariadne (Drosophila) homolog 2
        125310 RC_Z40646
                                R59161
                                             Hs.124953
                                                           ESTs
                                             Hs.106296
        125315 RC_Z41697
                                R38110
                                                           FSTs
15
                                                           ESTs, Wealthy similar to I38022 hypothetical protein [H.sapiens]
                                             Hs.112461
        125317 RC_Z99349
                                Z99348
                                AA081258
                                             Hs.132390
        135096 RC_Z99394_s
                                                           zinc finger protein 36 (KOX 18)
        104786 RC_AA027168
                                AA027167
                                             Hs.10031
                                                           KIAA0955 protein
                                                           EGF-TM7-latrophilin-related protein
        132837 D58024 s
                                AA370362
                                             Hs.57958
        120456 RC_AA251113
                                AA488750
                                             Hs.88414
                                                           BTB and CNC homology 1, basic leucine zipper transcription factor 2
20
                                                           fibronectin leucine rich transmembrane protein 2
        132459 RC_AA347573
                                             Hs.48998
                                AL120071
                                             Hs.154210
        101545 M31210
                                BE246154
                                                           endothelial differentiation, sphingolipid G-protein-coupled receptor, 1
        133505 C01527
                                AI630124
                                             Hs.324504
                                                           Homo sapiens mRNA; cDNA DKFZp586J0720 (from clone DKFZp586J0720)
        132360 RC N62948 s
                                AW893660
                                             Hs.46440
                                                           solute carrier family 21 (organic anion transporter), member 3
                                             Hs.264636
                                                           hypothetical protein FLJ20731
        132738 RC_W42674
                                AK000738
        119586 RC_W43000_s
129914 RC_N31750_s
25
                                AF088033
                                             Hs.159225
                                                           ESTs
                                NM_012421
                                             Hs.13321
                                                           rearranged L-myc fusion sequence
        130839 AF009301
                                AB011169
                                             Hs.20141
                                                           similar to S. cerevisiae SSM4
                                BE313625
                                             Hs.57435
                                                           solute carrier family 11 (proton-coupled divalent metal ion transporters), member 2
        132813 L37347
        134342 M99564
                                NM_000275
                                             Hs.82027
                                                           oculocutaneous albinism II (pink-eye dilution (murine) homolog)
30
                                AA083764
                                                           hypothetical protein MGC3178
        131878 RC_AA430673
                                             Hs.6101
        105426 RC_AA251297
                                W20027
                                              Hs.23439
                                                           ESTs
        132968 RC_AA620722
                                AF234532
                                             Hs.61638
                                                           myosin X
        132173 RC_W46577_s
                                X89426
                                             Hs.41716
                                                           endothellal cell-specific molecule 1
        113932 RC_W81237
114452 RC_AA020825
                                AA256444
                                                           hypothetical protein FLJ12604; KIAA1692 protein
                                             Hs.126485
                                                           Homo sapiens cDNA FLJ14445 fis, clone HEMBB1001294, highly similar to GTP-BINDING
35
                                Al369275
                                             Hs.243010
        PROTEIN TC10
                                AA806600
        115243 RC_AA278766
                                             Hs.116665
                                                           KIAA1842 protein
        134403 RC_H93708_s
                                AA334551
                                             Hs.82767
                                                           sperm specific antigen 2
        129647 RC_N49394
                                AB018259
                                             Hs.118140
                                                           KIAA0716 gene product
40
        111428 RC_H56559_s
                                AL031428
                                             Hs.174174
                                                           KIAA0601 protein
        115967 RC_AA446887
                                             Hs.42911
                                AI745379
                                                           ESTs
        120726 RC_AA293656
                                AA293655
                                             Hs.97293
                                                           ESTs
        114995 RC_AA251152
                                             Hs.193657
                                AA769266
                                                           ESTs
        303876 RC_AA233334_s U64820
                                             Hs.66521
                                                           Machado-Joseph disease (spinocerebellar ataxia 3, olivopontocerebellar ataxia 3, autosomal
45
        dominant, ataxin 3)
                                                           cytochrome b5 reductase b5R.2 ESTs
        311463 RC_Z39920
                                R55344
                                              Hs.22142
        120302 RC_AA192173
                                AA837098
                                             Hs.269933
        133071 RC_AA455044
                                BE384932
                                             Hs.64313
                                                           ESTs, Weakly similar to AF257182 1 G-protein-coupled receptor 48 [H.sapiens]
        121032 RC_AA398504
                                AA393037
                                             Hs.161798
                                                           ESTs
50
        129829 U41813
                                AF010258
                                              Hs.127428
                                                           homeo box A9
                                AW959615
        120245 RC_AA166965
                                             Hs.111045
                                                           ESTs
        120985 RC_AA398222
                                Al219896
                                             Hs.97592
                                                           ESTs
        114184 RC_Z39095
                                R56434
                                             Hs.21062
                                                           ESTs
                                                           Homo sapiens, Similar to RIKEN cDNA 1810038N03 gene, done MGC:9890, mRNA, complete
        447503 RC_AA284744_f AA115496
                                             Hs.336898
55
        132837 RC_AA428201
                                AA370362
                                             Hs.57958
                                                           EGF-TM7-latrophilin-related protein
        121034 RC_AA398507
                                AL389951
                                             Hs.271623
                                                           nucleoporin 50kD
        119718 RC_W69216
120455 RC_AA251083
                                W69216
                                             Hs.92848
                                                           ESTs
                                             Hs.104347
                                                           ESTS, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C WARNING ENTRY !!! [H.sapiens]
                                AA251720
60
                                              Hs.106932
        125280 RC_W93659
                                AI123705
                                                           ESTs
        132155 RC_AA227903
                                AK001607
                                             Hs.41127
                                                           hypothetical protein FLJ13220
        120609 RC_AA283902
                                AW978721
                                             Hs.266076
                                                           ESTs, Wealty similar to A46010 X-linked retinopathy protein [H.sapiens]
                                                           Homo sapiens cDNA FLJ11490 fis, clone HEMBA1001918
        121278 RC_AA401631
                                AA037121
                                             Hs.98518
        109023 RC_AA157293
                                AA157293
                                             Hs.72168
                                                           ESTs
        129815 RC_D60208_f
                                                           hypothetical protein FLJ21657
65
                                BE565817
                                             Hs.26498
        108061 RC_AA043979
                                AA043979
                                             Hs.62651
                                                           EST
        113287 RC_T66847
                                T66847
                                             Hs.194040
                                                           ESTs, Weakly similar to I38022 hypothetical protein [H.sapiens]
        114082 RC_Z38239
                                AK001612
                                             Hs.26962
                                                           Homo sapiens cDNA FLJ10750 fis, clone NT2RP3001929
        116334 RC_AA491457
                                                           ESTs
                                AL038450
                                             Hs.48948
70
        131486 RC_Z40071_s
                                F06972
                                              Hs.27372
                                                           BMX non-receptor tyrosine kinase
        107860 RC_AA024961
                                AA024961
                                             Hs 50730
                                                           FSTs
                                AU077002
                                             Hs.24950
                                                           regulator of G-protein signalling 5
        131263 RC_AA443826
        132207 RC_AA443294
                                BE206939
                                             Hs.42287
                                                           E2F transcription factor 6
        129183 RC_AA155743
                                BE561824
                                              Hs.273369
                                                           uncharacterized hematopoietic stem/progenitor cells protein MDS027
75
                                AJ338631
                                              Hs.43266
                                                           Homo sapiens cDNA: FLJ22536 fis. clone HRC13155
        408431 RC_T23708
                                AW978022
                                             Hs.238911
                                                           hypothetical protein DKFZp762E1511; KIAA1816 protein
        120575 RC_AA280934
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132121 RC_AA443284_s NM_004529
                                             Hs.404
                                                           myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog); translocated to, 3
        117657 RC N39074
                                N39074
                                              Hs.44933
                                                           ESTs
                                                           prefoldin 4
        134922 RC_W04507_s
                                AI718295
                                              Hs.91161
                                                           myosin VA (neavy polypeptide 12, myoxin)
gb:ns44f05.s1 NCI_CGAP_Alv1 Homo sapiens cDNA clone, mRNA sequence
        118523 RC_R41828_s
                                Y07759
                                              Hs.170157
 5
         116845 RC_H64973
                                AA649530
         115291 RC_AA279943
                                BE545072
                                              Hs.122579
                                                           hypothetical protein FLJ10461
         120326 RC_AA196300
                                AA196300
                                              Hs.21145
                                                           hypothetical protein RG083M05.2
                                                           protein phosphatase 3 (formerly 2B), catalytic subunit, beta isoform (calcineurin A beta)
         130174 M29550
                                M29551
                                              Hs.151531
         129131 RC_AA436489
                                                           dual specificity phosphatase 10
                                AB026436
                                              Hs.177534
10
         129868
                RC_AA287032
                                AW172431
                                              Hs.13012
                                                           ESTs
         118661 RC_N70777
                                AL137554
                                              Hs.49927
                                                           protein kinase NYD-SP15
         129829 RC_AA496921
                                AF010258
                                              Hs.127428
                                                           homeo box A9
                                                           ESTs, Wealty similar to T08599 probable transcription factor CA150 [H.sapiens]
        115985 RC_AA447709
                                              Hs.268115
                                AA447709
        134637 RC_AA369856_s
                                U87309
                                              Hs.180941
                                                           vacuolar protein sorting 41 (yeast homolog)
15
         132714 RC_AA252598
                                W39388
                                              Hs.55336
                                                           Homo sapiens, clone MGC:17421, mRNA, complete cds
         129771 RC_H73237
                                              Hs.102708
                                                           DKFZP434A043 protein
                                AL096748
         123360 RC_AA504784
                                AA532718
                                              Hs.178604
                                                           ESTs
                                                           hypothetical protein FLJ10808
        132902 RC_AA490969
                                A1936442
                                              Hs.59838
        113716 RC_T97750
113825 RC_W48860
                                AA001356
                                              Hs.18159
                                                           ESTs
20
                                AW014486
                                              Hs.22509
                                                           ESTs
         130367 RC_Z38501
                                AL135301
                                              Hs.8768
                                                           hypothetical protein FLJ10849
                                W07318
                                                           M-phase phosphoprotein 1
         120541 RC_AA278298
                                              Hs.240
        116727 RC F13684
                                R76472
                                              Hs.65646
                                                           ESTs
                                AA862391
                                              Hs.48494
                                                           ESTs, Moderately similar to A46010 X-linked retinopathy protein [H.sapiens]
         118219 RC_N62231
25
         119767 RC_W72562
                                W72562
                                              Hs.58119
                                                           ESTs
                                                           oncogene TC21
         128917 RC_AA481252
                                Al365215
                                              Hs.206097
        451553 RC_AA020928
132716 RC_AA251288
                                AA018454
                                              Hs.269211
                                                           ESTs
                                BE379595
                                              Hs.283738
                                                           casein kinase 1, atpha 1
         118525 RC_N67861
                                N67861
                                              Hs.49390
                                                           ESTs
30
         114618 RC_AA084162
                                AW979261
                                              Hs.291993
                                                           ESTs
         119743 RC_W70242
                                AA947552
                                              Hs.58086
                                                           ESTs
         108154 RC_AA425151_s NM_005754
                                              Hs.220689
                                                           Ras-GTPase-activating protein SH3-domain-binding protein
         122798 RC_AA460324
                                AW366286
                                              Hs.145696
                                                           splicing factor (CC1.3)
                                AW410035
                                              Hs.75862
                                                            MAD (mothers against decapentaplegic, Drosophila) homolog 4
         133746 U44378
35
         119822 RC_W74471
                                AF086409
                                              Hs.301327
                                                           FSTs
         122186 RC_AA435842
                                                           ESTs
                                AA398811
                                              Hs.104673
         114941 RC_AA243017
                                AA236512
                                              Hs.87331
                                                            ESTs
         118053 RC_N53367
                                N53391
                                              Hs.47629
                                                            ESTs
                                                           down-regulator of transcription 1, TBP-binding (negative cofactor 2)
         123234 RC_AA490227
                                NM_001938
                                              Hs.16697
40
                                                           gastric intrinsic factor (vitamin B synthesis)
        129280 M63154
                                M63154
                                              Hs.110014
         118995 RC_N94591
                                N94591
                                                           ESTs
                                              Hs.323056
         116750 RC_H05960
                                AA760689
                                              Hs.92418
                                                            ESTs
         129026 M98833
                                AL120297
                                              Hs.108043
                                                            Friend leukemia virus integration 1
         105127 RC_AA158132
                                AA045648
                                              Hs.301957
                                                            nudix (nucleoside diphosphate linked moiety X)-type motif 5
45
                                              Hs.103446
         114513 RC_AA044825
                                AA044873
                                                            ESTs
                                                            Homo saniens cDNA: FLJ23269 fis, done COL09533
        411856 RC_T35697
                                H67899
                                              Hs.4190
                                                           hypothetical protein DKFZp434E2220
         132036 W01568
                                AL157433
                                              Hs.37706
         130091 RC_W88999
                                W88999
                                                           gb:zh70h03.s1 Soares_fetal_fiver_spleen_1NFLS_S1 Homo sapiens cDNA clone 3', mRNA
         sequence
50
         414108 U09564
                                Al267592
                                              Hs.75761
                                                            SFRS protein kinase 1
         119881 RC_W81456
                                W81486
                                              Hs.58648
                                                           ESTs
        117770 RC_N47953
                                AW957372
                                              Hs.46791
                                                           ESTs, Weakly similar to 138022 hypothetical protein [H.sapiens]
                                              Hs.58452
         119850 RC_W80447
                                Al247568
                                                           ESTs
         115439 RC_AA284561
                                AI567972
                                              Hs.193090
                                                           ESTs, Highly similar to AF161437 1 HSPC319 [H.sapiens]
55
         123107 RC_AA486071
                                AA225048
                                              Hs.104207
         406698 M24364
                                X03068
                                              Hs.73931
                                                            major histocompatibility complex, class II, DQ beta 1
         121231 RC_AA400780
                                AA814948
                                                           ESTs, Weakly similar to ALUC_HUMAN !!!! ALU CLASS C WARNING ENTRY !!! [H.sapiens]
                                              Hs.96343
                                AA478486
                                              Hs.3852
         132074 AB002366
                                                            KIAA0368 protein
                                AB000115
         413670 AB000115
                                              Hs.75470
                                                           hypothetical protein, expressed in osteoblast
60
         125277 RC_W93227
                                W93227
                                              Hs.103245
         114056 RC_AA186324
                                AA188175
                                              Hs.82506
                                                            KIAA1254 protein
         121153 RC_AA399640
                                AA399640
                                              Hs.97694
                                                           ESTs
         121609 RC_AA416867
                                AA416867
                                              Hs.98185
                                                            EST
                                                            ESTs, Weakly similar to ALUB_HUMAN !!!! ALU CLASS B WARNING ENTRY !!! [H.sapiens]
                                AA287556
         120661 RC_AA287556
                                              Hs.263412
65
         120850 RC_AA349647
                                AA349647
                                              Hs.96927
                                                            Homo sapiens cDNA FLJ12573 fis, clone NT2RM4000979
         124947 RC_T03170
                                T03170
                                              Hs.100165
         130529 RC_AA280886
                                AA178953
                                                           gb:zp39e03.s1 Stratagene muscle 937209 Homo sapiens cDNA clone 3' similar to contains Alu
         repetitive element, mRNA sequence
         117683 RC_N40180
                                                            gb:yy44d02.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA clone
                                N40180
70
         IMAGE:276387 3' similar to contains L1.t1 L1 repetitive element; mRNA sequence.
                                                            gb:EST10426 Adipose tissue, white I Homo sapiens cDNA 3' end, mRNA sequence.
         120745 RC_AA302809
                                AA302809
         120936 RC_AA385934
                                AA385934
                                              Hs.97184
                                                            EST, Highly similar to (defilne not available 7499603) [C.elegans]
         112597 RC_R78376
                                R78376
                                              Hs.29733
                                                            EST
         120183 RC_Z40174
                                AW082866
                                              Hs.65882
                                                           ESTs
75
         120644 RC_AA287038
                                AJ869129
                                              Hs.96616
                                                            ESTs
```

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N98488
                                                           gb:zb82h01.s1 Soares_senescent_fibroblasts_NbHSF Homo sapiens cDNA clone
        119023 RC_N98488
        IMAGE:310129 3', mRNA sequence.
                                             Hs.59952
        107582 RC_AA002147
                                AA002147
                                N62580
                                             Hs.322925
                                                           EST, Weakly similar to putative p150 [H.sapiens]
        118249 RC_N62580
        115022 RC_AA252029
                                AA252029
                                             Hs.87935
                                                           ESTs
        117710 RC_N45198
                                N45198
                                             Hs.47248
                                                           ESTs. Highly similar to similar to Cdc14B1 phosphatase [H.sapiens]
        115341 RC_AA281452
                                AA281452
                                             Hs.88840
                                                           EST, Weakly similar to granute cell marker protein [M.muscutus]
        118896 RC_N90680
                                N46213
                                             Hs.54642
                                                           methionine adenosyltransferase II, beta
        121121 RC_AA399371
                                AA399371
                                             Hs.189095
                                                           similar to SALL1 (sal (Drosophila)-like
10
        118329 RC_N63520
                                N63520
                                                           gb:yy62f01.s1 Soares_multiple_sclerosis_2NbHMSP Homo saptens cDNA done IMAGE:278137
        3', mRNA sequence.
        119496 RC_W35416
118111 RC_N55493
                                                           ESTs, Moderately similar to A46010 X-linked retinopathy protein [H.saplens]
                                W35416
                                             Hs.156861
                                                           gb:yv50c02.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:246146 3',
                                N55493
        mRNA sequence.
15
        119062 RC_R16698
                                AW444881
                                             Hs.77829
        116710 RC_F10577_f
                                                           v-crk avian sarcoma virus CT10 oncogene homolog
                                F10577
                                             Hs.306088
        119261 RC_T15956
                                T15956
                                             Hs.65289
                                                           gb:aa86b10.s1 Stratagene fetal retina 937202 Homo sapiens cDNA clone IMAGE:838171 3'
        122723 RC_AA457380
                                AA457380
        similar to contains L1.b3 L1 repetitive element;, mRNA sequence.
20
        117732 RC_N46452
                                N46452
                                                           gb:yy76h09.s1 Soares_multiple_sclerosis_2NbHMSP Homo sapiens cDNA done
        IMAGE:279521 3' similar to contains L1.t2 L1 repetitive element; mRNA sequence.
                                                           gbrze97d11.s1 Soares_fetal_heart_NbHH19W Homo sapiens cDNA clone IMAGE:366933 3"
         104787 RC_AA027317
                                AA027317
        similar to contains Alu repetitive element;, mRNA sequence.
        100071 A28102
                                A28102
                                                           Human GABAa receptor alpha-3 subunit
25
        115819 RC_AA426573
                                AA486620
                                             Hs.41135
                                                           endomucin-2
        130882 RC Z40166 f
                                AA497044
                                             Hs.20887
                                                           hypothetical protein FLJ10392
        125225 RC_W76540
108339 RC_AA070801
                                W74169
                                             Hs.16492
                                                           DKFZP564G2022 protein
                                                           ESTS, Weakly similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE CONTAMINATION
                                AW151340
                                             Hs.51615
        WARNING ENTRY [H.sapiens]
30
        100338 D63483
                                D86864
                                             Hs.57735
                                                           acetyl LDL receptor; SREC
        121636 RC_AA417027
                                AA379203
                                             Hs.306654
                                                           Homo sapiens cDNA FLJ13574 fis, clone PLACE1008625
                                T26379
                                                           Homo sapiens done 23632 mRNA sequence
         103875 RC_AA418387
                                             Hs.48802
        118716 RC_N73460
                                                           fucosyltransferase 8 (alpha (1,6) fucosyltransferase)
                                AI658908
                                             Hs.118722
                                                           Homo sapiens cDNA: FLJ22063 fis, clone HEP10326
         119763 RC_W72450
                                R54146
                                             Hs.10450
35
        121917 RC_AA428218
                                AA406397
                                             Hs.98038
                                                           ESTs
        132806 M91488
                                A1699432
                                              Hs.278619
                                                           hypothetical protein FLJ10099
                                             Hs.285115
                                                           interleukin 13 receptor, alpha 1
         130949 Y10659
                                AV656840
         108806 RC_AA129933
                                AF070578
                                             Hs.71168
                                                           Homo sapiens done 24674 mRNA sequence
         133276 RC_AA490478
                                             Hs.69504
                                AW978439
                                                           ESTs
40
                                             Hs.89548
                                NM_000121
                                                           erythropaietin receptor
         134760 RC_H16758
         132867 AA121287
                                AF226667
                                             Hs.58553
                                                           CTP synthase II
                                AA393968
                                             Hs.180145
                                                           HSPC030 protein
         132051 AA091284
        114208 RC_Z39301
                                AL049466
                                             Hs.7859
                                                           ESTs
         104094 AA418187
                                AA418187
                                              Hs.330515
                                                           ESTs
45
                                NM_002959
         128718 AA426361
                                             Hs.281706
                                                           sortilin 1
                                NM_001992
                                             Hs.128087
                                                           coagulation factor II (thrombin) receptor
         302032 RC_N20407
         115501 RC_AA291553
                                AA291553
                                             Hs.190086
                                                           ESTs
                                AU076536
                                              Hs.50984
         101997 U01160
                                                           sarcoma amplified sequence
                                                           hypothetical protein FLJ20038
                                AA430591
                                              Hs.72071
         103708 AA037206
50
                                                           RYK receptor-like tyrosine kinase
         101899 $59184
                                $59184
                                              Hs.79350
                                             Hs.28935
                                                           transducin-like enhancer of split 1, homolog of Drosophila E(sp1)
         115839 RC_AA429038
                                BE300266
                                              Hs.54481
                                                           low density lipoprotein receptor-related protein 8, apolipoprotein e receptor
         409459 D50678
                                D86407
                                              Hs.150402
                                                           Activin A receptor, type I (ACVR1) (ALK-2)
         103563 Z22534
                                L02911
                                                           ESTs, Weakly similar to MAPB_HUMAN MICROTUBULE-ASSOCIATED PROTEIN 1B
                                AW974175
                                             Hs.188751
         123233 RC_AA490225
55
        (H.sadens)
         121305 RC_AA402468
                                AA402468
                                              Hs.291557
                                              Hs.54900
                                                           serologically defined colon cancer antigen 1
         114798 RC_AA159181
                                AA159181
                                                           Homo sapiens, clone IMAGE:2961368, mRNA, partial cds
                                              Hs.6592
         133145 RC_AA196549
                                H94227
         131567 RC_AA291015_s AF015592
                                              Hs.28853
                                                           CDC7 (cell division cycle 7, S. cerevisiae, homolog)-like 1
60
        112300 RC_R54554
                                H24334
                                              Hs.26125
         129507 RC_AA192099
                                AJ236885
                                              Hs.112180
                                                           zinc finger protein 148 (pHZ-52)
         121033 RC_AA398505
                                AA398505
                                              Hs.97360
                                                           ESTs
                                              Hs.143629
         121151 RC_AA399636
                                AA399636
                                                           ESTs
         121402 RC_AA406063
                                AA406063
                                              Hs.98003
                                                           ESTs
65
         123203 RC_AA489671
                                AA352335
                                              Hs.65641
                                                           hypothetical protein FLJ20073
                                AB030034
                                              Hs.115175
                                                           sterile-alpha motif and leucine zipper containing kinase AZK
         132271 RC_AA236466
         125197 RC_W69106
                                AF086270
                                              Hs.278554
                                                           heterochromatin-like protein 1
                                              Hs.290880
                                                           ESTs, Wealty similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION
         114935 RC_AA242809
                                H23329
         WARNING ENTRY [H.sapiens]
70
                                AW401809
         125279 RC_W93640
                                              Hs.4779
                                                           KIAA1150 protein
         108778 RC_AA128548
                                AF133123
                                              Hs.90847
                                                           general transcription factor IIIC, polypeptide 3 (102kD)
                                AA045708
                                              Hs.40545
         108087 RC AA045709
                                                           ESTs
         132466 RC_N66810_s
                                AJ597655
                                              Hs.49265
                                                           ESTs
         133328 R36553
                                AW452738
                                              Hs.265327
                                                           hypothetical protein DKFZp761I141
                                              Hs.73853
75
         124057 RC_F13604
                                AA902384
                                                           bane morphogenetic protein 2
         124800 RC_R45115
                                AW864086
                                              Hs.138617
                                                           thyroid hormone receptor interactor 12
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121029 RC_AA398482
                                AA398482
                                             Hs.97641
                                                          EST
        120663 RC_AA287627
                                AA827798
                                             Hs.105089
                                                          ESTs
                                             Hs.155596
                                                          BCL2/adenovirus E1B 19kD-interacting protein 2
        102133 U15173
                                AU076845
                                             Hs.146343
        108246 RC_AA062855
                                AH23132
                                                          ESTs
 5
                                                           N-myristoyttransferase 2
                                             Hs.122647
        125226 RC_W78134
                                AA782536
        120260 RC_AA171739
                                AK000061
                                             Hs.101590
                                                          hypothetical protein
                                             Hs.107815
        124906 RC_R87647
                                H75964
                                                           ESTs
        109406 RC_AA226877
                                AA199883
                                             Hs.67624
                                                          ESTs
        109271 RC_AA195668
                                AW137422
                                             Hs.86022
                                                          ESTs
10
                                             Hs.222779
                                                           ESTs, Moderately similar to similar to NEDD-4 [H.sapiens]
        125052 RC_T80174_s
                                T85104
        109101 RC_AA167708
                                AW608930
                                             Hs.52184
                                                           hypothetical protein FLJ20618
        115241 RC_AA278723
                                AA648278
                                             Hs.193859
                                                           ESTs
        117163 RC_H97909
                                N36861
                                             Hs.42344
                                                           ESTs
                                                           ESTs
                                             Hs.16732
        113530 RC_T90313
                                T90313
15
                                                           Zic family member 3 (odd-paired Drosophila homolog, heterotaxy 1)
        120375 RC_AA227260
                                AF028706
                                             Hs.111227
        129435 AA314256
                                AF151852
                                             Hs.111449
                                                           CGI-94 protein
        114864 RC_AA235256
                                AA135332
                                             Hs.71608
                                                           ESTs
                                                           ADP-ribosytation factor-like 5
        103988 AA314389
                                AA314389
                                             Hs.42500
                                                           CDC14 (cell division cycle 14, S. cerevisiae) homolog B
                                             Hs.22116
        131006 RC_AA242763
                                AF064104
20
        106781 RC_AA478474
                                AA330310
                                             Hs.24181
                                                           ESTs
        106141 RC_AA424558
                                AF031463
                                             Hs.9302
                                                           phosducin-like
                                                           hypothetical protein MGC10947
        116213 RC_AA476738
                                AA292105
                                             Hs.326740
                                             Hs.97393
                                                           KIAA0328 protein
        135266 AB002326
                                 R41179
        135058 RC_AA430152
                                AJ379720
                                             Hs.93814
                                                           hypothetical protein
25
        119908 RC_W85844
                                 AA524470
                                             Hs.58753
                                                           ESTs
        103695 AA018758
                                 AW207152
                                             Hs.186600
                                                           ESTs
                                                           chromosome 21 open reading frame 6
        103978 AA307443
                                 NM_016940
                                             Hs.34136
         109485 RC_AA233472
                                BE619092
                                             Hs.28465
                                                           Homo sapiens cDNA: FLJ21869 fis, clone HEP02442
        129574 AA458603
                                 AA026815
                                             Hs.11463
                                                           UMP-CMP kinase
30
        115347 RC_AA281528
                                 AA356792
                                             Hs.334824
                                                           hypothetical protein FLJ14825
                                                           ESTs, Weakly similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION
                                 AW961026
                                             Hs.96752
         120765 RC_AA338735
        WARNING ENTRY [H.sapiens]
         121059 RC_AA398628
                                AA393283
                                                           gb:zt74e03.r1 Soares_testis_NHT Homo sapiens cDNA clone 5', mRNA sequence
         131887 AA046548 ·
                                             Hs.332848
                                                           SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily e,
                                 W17064
35
        member 1
                                                           Homo sapiens mRNA; cDNA DKFZp586O1318 (from clone DKFZp586O1318)
                                             Hs.22689
         112064 RC_R43812
                                 AL049390
         115606 RC_AA400465
                                 AI025829
                                             Hs.86320
                                                           core-binding factor, runt domain, alpha subunit 2; translocated to, 1; cyclin D-related
         131750 RC_H94855_s
                                 NM_004349
                                             Hs.31551
                                             Hs.1594
         102123 U14518
                                 NM_001809
                                                           centromere protein A (17kD)
         129847 RC_W46767
133809 RC_AA235275
40
                                                           hypothetical protein FLJ22637
                                 N64025
                                              Hs.296178
                                 AV649326
                                             Hs.76359
                                                           catalase
                                 NM_007203
         132210 RC_N51499_s
                                             Hs.42322
                                                           A kinase (PRKA) anchor protein 2
                                             Hs.98390
         122356 RC_AA443794
                                 AA443794
                                                           ESTs
                                             Hs.42369
         114958 RC_AA243708
                                 N20912
                                                           ESTs
45
                                                           Homo sapiens mRNA; cDNA DKFZp761J1112 (from clone DKFZp761J1112)
         103951 AA287840
                                 AL353944
                                             Hs.50115
         134703 RC_AA280704
                                 AF117065
                                             Hs.88764
                                                           male-specific lethal-3 (Drosophila)-like 1
         128727 AA287864
                                 A1223335
                                              Hs.50651
                                                           Janus kinase 1 (a protein tyrosine kinase)
                                                           sema domain, immunoglobulin domain (Ig), transmembrane domain (TM) and short cytoplasmic
         105743 RC_AA293300_s BE246502
                                             Hs.9598
         domain, (semaphorin) 4B
50
         103744 AA076003
                                 AA079267
                                                           gb:zm97e10.s1 Stratagene colon HT29 (937221) Homo sapiens cDNA clone 3', mRNA
         sequence
         114348 N80402
                                 AL050321
                                             Hs.301532
                                                           CRP2 binding protein
                                                           KIAA0831 protein
         114009 RC_W90067
                                 Al248544
                                             Hs.103000
                                 AA837124
         134704 RC_AA280849
                                              Hs.88780
                                                           EST<sub>S</sub>
55
                                                           DKFZP434A043 protein
         128629 AA399187
                                 AL096748
                                              Hs.102708
                                                           Homo saplens cDNA FLJ13694 fis, clone PLACE2000115
         104410 H65925
                                 AI807519
                                              Hs.104520
                                              Hs.31802
                                                           ESTs, Highly similar to A59266 unconventional myosin-15 [H.sapiens]
         110200 RC_H21075
                                 H21075
         124483 RC_N53976
                                 AI821780
                                              Hs.179864
         101391 M14648
                                 NM_002210
                                              Hs.295726
                                                           integrin, alpha V (vitronectin receptor, alpha polypeptide, antigen CD51)
60
         109657 RC_F04826
                                              Hs.26814
                                                           ESTs
                                 R60900
         117140 RC_H96813
                                 H96813
                                              Hs.42241
                                                           ESTs
         132937 RC_AA233706_f
                                 AW952912
                                              Hs.300383
                                                           hypothetical protein MGC3032
                                              Hs.239114
         129799 R36410
                                 AW967473
                                                           mannosidase, alpha, class 1A, member 2
         105077 RC_AA142919
                                                           Homo sapiens cDNA FLJ12082 fis, done HEMBB1002492
                                              Hs.234863
                                 W55946
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                                 AA836472
                                              Hs.297939
                                                           cathensin B
         100850 RC_N58561_s
                                                           epilepsy, progressive myodonus type 2, Lafora disease (laforin) gb:Human endogenous retrovirus K clone 10.1 polymerase mRNA, partial cds
         131043 RC_AA490925
                                 AF084535
                                              Hs.22464
         118417 RC_N66048_f
                                 AF080229
         129254 RC_AA243695
                                 AA252468
                                              Hs.1098
                                                           DKFZp434J1813 protein
                                              Hs.65732
         119149 RC_R58910
                                 BE304701
                                                           ESTs
                                                           DKFZP434F2021 protein
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         133996 AA091367
                                 AA380267
                                              Hs.78277
         110223 RC_H23747
                                              Hs.31697
                                 H19836
                                                           EST<sub>8</sub>
                                              Hs.281348
                                                           hypothetical protein FLJ10895
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                                 AK001757
                                              Hs.97849
                                                           ESTs
         135286 RC_AA424469_s
                                 AW023482
                                 AA806187
                                              Hs.289101
                                                           glucose regulated protein, 58kD
         122967 RC_AA478521
75
                                                            ubiquitination factor E4B (homologous to yeast UFD2)
         131236 AA282640
                                 AF043117
                                              Hs.24594
         128568 AA463380
                                 H12912
                                              Hs.274691
                                                           adenytate kinase 3
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112888 RC_T03872
                                AW195317
                                             Hs.107716
                                                          hypothetical protein FLJ22344
        115192 RC_AA261920
                                AA741024
                                             Hs.88378
                                                          ESTs
                                                          hypothetical protein FLJ20701
        118688 RC_N71484
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                                             Hs.169764
                                                          gbrzv57g07.s1 Soares_testis_NHT Homo sapiens cDNA clone 3', mRNA sequence
        122264 RC_AA436837
                                AA436837
                                                          CGG triplet repeat binding protein 1
        128981 AA135452
                                AA927177
                                             Hs.86041
                                                          hypothetical protein MGC2628
        131042 RC_R42457
                                A1826288
                                             Hs.171637
                                                          hypothetical protein FLJ21062
                                             Hs.151258
        103704 AA028171
                                AA028171
                                                          MAD (mothers against decapentaplegic, Drosophila) homolog 6
        121341 AA233107
                                AF035528
                                             Hs.153863
        106593 RC_AA456826
                                AW296451
                                             Hs.24605
        115195 RC_AA262156
115425 RC_AA284071
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                                AW968619
                                             Hs.155849
                                                          ESTs
                                             Hs.180680
                                                          ESTs. Weakly similar to 154374 gene NF2 protein [H.sapiens]
                                AA811895
        117258 RC_N21299
                                AF086041
                                             Hs.42975
                                                          ESTs
        120209 RC_Z40892
                                F02951
                                                          gb:HSC1HB082 normalized infant brain cDNA Homo sapiens cDNA clone c-1hb08 3', mRNA
        sequence
15
         134082 L16991
                                L16991
                                             Hs.79006
                                                          deoxythymidylate kinase (thymidylate kinase)
        104774 RC_AA026066
                                AW959755
                                             Hs.288896
                                                          Homo sapiens cDNA FLJ12977 fis, done NT2RP2006261
        115625 RC_AA401630
                                             Hs.62592
                                AA059459
                                                          Homo sapiens chromosome 19, BAC 282485 (CIT-B-344H19)
        104469 N28707
                                N28707
                                             Hs.154304
        107401 W20054
                                N91453
                                             Hs.102987
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        111686 RC_R21510
                                R22039
                                             Hs.23217
                                                          ESTs
        115300 RC_AA280026
                                AA280095
                                             Hs.88689
                                                          ESTs
                                                          hypothetical protein FLJ10335
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                                AA282292
                                             Hs.279841
                                N41549
                                             Hs.285410
        132224 RC_H97819
                                                          ESTs
        113791 M95767
                                Al269096
                                             Hs.135578
                                                          chitobiase, di-N-acetyl-
25
                                                          hypothetical protein DKFZp434P0116
         129144 AA004987
                                             Hs.20137
                                AL137275
        104448 L44574
                                NM 007331
                                             Hs.110457
                                                          Wolf-Hirschhorn syndrome candidate 1
                                NM_002267
                                                          karyopherin alpha 3 (importin alpha 4)
         132084 RC_T26981_s
                                             Hs.3886
        111831 RC_R36083
                                             Hs.268695
                                R36095
                                                          ESTs
                                             Hs.337532
                                                          ESTs, Weakly similar to A47582 B-cell growth factor precursor [H.sapiens]
        114765 RC_AA252163
                                AA463550
30
        115029 RC_AA252219
                                AL137939
                                             Hs.40096
        100457 H81492
                                BE246400
                                             Hs.285176
                                                          acetyl-Coenzyme A transporter
                                                           Homo sapiens cDNA FLJ14673 fis, clone NT2RP2003714, moderately similar to ZINC FINGER
         104536 R24011
                                R24024
                                             Hs.158101
        PROTEIN 91
                                AI091731
         116167 RC_AA461562
                                             Hs 87293
                                                           hypothetical protein FLJ20045
35
        103889 AA236771
                                R85350
                                             Hs.101368
                                                          ESTs
        131978 RC_H48459_s
                                AA355925
                                             Hs.36232
                                                           KIAAD186 gene product
                                N80181
                                              Hs.221498
        118843 RC_N80181
                                                          ESTs
         120837 RC W93092
                                                          Homo sapiens cDNA FLJ11963 ffs, clone HEMBB1001051
                                BE149656
                                             Hs.306621
                                                           KIAA0029 protein
                                NM_015361
                                             Hs 268053
         133647 D21852
40
                                                          nucleoporin 98kD
         129521 U41815
                                AF071076
                                             Hs.112255
                                                           gb:zm83c07.s1 Stratagene ovarian cancer (937219) Homo sapiens cDNA clone 3', mRNA
         103746 AA081876
                                AA075000
        sequence
         132019 RC_AA134965_i H56995
                                             Hs,37372
                                                           Homo saplens DNA binding peptide mRNA, partial cds
         132310 RC_AA284107
                                             Hs.289044
                                                           Homo sapiens cDNA FLJ12048 fis, done HEMBB1001990
                                AA173223
45
         117367 RC_N24954
                                             Hs.42502
                                AI041793
                                                           ESTs
                                                           gb:zm89b09.r1 Stratagene ovarian cancer (937219) Homo sapiens cDNA clone 5' similar to
         103743 AA075998
                                AA075998
        gb:M15887 ACYL-COA-BINDING PROTEIN (HUMAN);, mRNA sequence
                                                          gb:nz79b10.s1 NCL_CGAP_GCB1 Homo sapiens cDNA clone 3' similar to gb:M34539 FK506-
         103761 AA085138
                                AA765163
        BINDING PROTEIN (HUMAN);, mRNA sequence
50
                                AA913909
                                             Hs.153088
                                                          TATA box binding protein (TBP)-associated factor, RNA polymerase I, A, 48kD
        130237 L39060
                                AA504428
                                             Hs.10487
         128752 RC_N72879
                                                           Homo sapiens, clone IMAGE:3954132, mRNA, partial cds
                                Al187925
                                             Hs.95667
         135162 AA045930
                                                          F-box protein 30
                                                           dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 2
        131386 AA096412
                                BE219898
                                             Hs.173135
        129021 RC_AA599244
                                AL044675
                                             Hs.173081
                                                           KIAA0530 protein
55
        424274 AA293634
                                W73933
                                             Hs.283738
                                                          caseln kinase 1, alpha 1
                                                           cAMP responsive element binding protein-like 2
         129913 H06583
                                NM_001310
                                             Hs.13313
                                                           Homo sapiens cDNA: FLJ22488 fis, clone HRC10948, highly similar to HSU79298 Human clone
        131888 U79298
                                AW294659
                                             Hs.34054
        23803 mRNA
        118612 RC_N69466
                                AB037788
                                             Hs.224961
                                                           cleavage and polyadenylation specific factor 2, 100kD subunit
60
        322026 AA203138
                                AW024973
                                             Hs.283675
                                                           NPD009 protein
        110892 RC N38882
                                AL035301
                                             Hs.97375
                                                           H.sapiens gene from PAC 106H8
                                                           ESTs, Wealty similar to 154374 gene NF2 protein [H.saptens]
        111429 RC_R01245
                                A1038052
                                             Hs.19162
        113334 RC_T76962
                                AW974666
                                             Hs.293024
                                                          FSTs
         104091 AA417310
                                BE465093
                                             Hs.106101
                                                           hypothetical protein FLJ22557
65
         105246 RC_AA226879
                                AA226879
                                                           gb:zr19c09.s1 Stratagene NT2 neuronal precursor 937230 Homo sapiens cDNA clone
        IMAGE:663856 3' similar to contains Alu repetitive elemen
                                                          t, mRNA sequence.
        113300 RC_T67448
                                T67448
                                             Hs.13101
                                                           ESTs
         117147 RC_H97225_s
                                AW901347
                                             Hs.38592
                                                           hypothetical protein FLJ23342
         121349 RC_AA405205
                                AA405205
                                             Hs.97960
                                                           ESTs, Weakly similar to T51146 ring-box protein 1 [H.sapiens]
70
         100294 D49396
                                AA331881
                                             Hs.75454
                                                           peroxiredoxin 3
         133999 M28213
                                AA535244
                                             Hs.78305
                                                           RAB2, member RAS oncogene family
                                                           Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 2004403
         133259 AA278548
                                BE379646
                                             Hs.6904
         129423 AA371418
                                AA204686
                                              Hs.234149
                                                           hypothetical protein FLJ20647
                                              Hs.236642
         131098 RC_AA459668
                                11666669
                                                           3-hydroxyisobutyryl-Coenzyme A hydrolase
75
                                 AI828337
         135272 AA399391
                                              Hs.97591
                                                           ESTs
                                                           Homo sapiens mRNA; cDNA DKFZp434P228 (from clone DKFZp434P228)
         129155 AA046865
                                AI952677
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311291 AA056319
                                 AA782601
                                              Hs.319817
        120750 RC_AA310499
                                 AJ191410
                                              Hs.96693
                                                            ESTs, Moderately similar to 2109260A B cell growth factor [H.saplens]
                                 AV655843
                                              Hs.169919
        101002 J04058
                                                            electron-transfer-flavoprotein, alpha polypeptide (glutaric aciduria II)
                                                            Homo sapiens, clone IMAGE:3351295, mRNA
        133012 AA099241
                                 AA847843
                                              Hs.62711
                                                            mitochondrial ribosomal protein L32
        103879 AA228148_s
                                 BE543269
                                              Hs.50252
        131281 RC_AA443212
                                 AA251716
                                              Hs.25227
        115109 RC_AA256383
                                                            protein kinase, AMP-activated, gamma 3 non-catalytic subunit
                                 AJ249977
                                              Hs.88049
                                                            Homo sapiens mRNA; cDNA DKFZp564B182 (from clone DKFZp564B182)
        118502 RC_N67317
                                 AL157488
                                              Hs.50150
        134100 L07540
                                 AA460085
                                              Hs.171075
                                                            replication factor C (activator 1) 5 (36.5kD)
10
        131869 AA484944
                                 AW958547
                                              Hs.33540
                                                            ESTs, Weakly similar to dJ309K20.4 [H.sapiens]
        115396 RC_AA282985
                                              Hs.89081
                                                            ESTs
                                 AA810854
                                 AW976877
        103860 AA203742
                                              Hs.38057
                                                            ESTs
        135089 N75611_s
                                 A1918035
                                              Hs.301198
                                                            roundabout (axon guidance receptor, Drosophila) homolog 1
         129938 U79300
                                 AW003668
                                               Hs.135587
                                                            Human clone 23629 mRNA sequence
15
                                                            Homo sapiens cDNA: FLJ21564 fis, clone COL06452
         107508 W90095
                                 N74925
                                               Hs.38761
        103685 AA005190
                                 AA158008
                                              Hs.292444
                                                            ESTs
                                 AL020996
                                               Hs.8518
         125170 AA203147
                                                            selenoprotein N
        129179 RC_AA504125_s
                                 AW969025
                                              Hs.109154
                                                            ESTs
        116262 AA477046
                                 A1936442
                                               Hs.59838
                                                            hypothetical protein FLJ10808
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                                               Hs.78305
                                                            RAB2, member RAS oncogene family
        123009 RC_AA479949
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        131004 D29833
                                 D29833
                                               Hs.2207
                                                            salivary proline-rich protein
                                               Hs.166091
                                                            ligase IV, DNA, ATP-dependent
         103317 X83441
                                 X83441
         132814 RC_C15251_f
                                 D60730
                                               Hs.57471
                                                            ESTs
         103992 U77718
                                 BE018142
                                               Hs.300954
                                                            Huntingtin interacting protein K
25
                                               Hs.84928
                                                            nuclear transcription factor Y, beta
         109258 X59710
                                 AL044818
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                                 AW302200
                                               Hs.6336
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         132727 AA136382_s
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                                 N27495
                                               Hs.5565
                                 AF032922
                                               Hs.8813
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         134664 AA256106
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                                               Hs.87507
                                                            ESTs
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         103826 AA165564
                                 AW162998
                                               Hs.24684
                                                            KIAA1376 protein
                                               Hs.169927
                                                            ESTs
        111678 RC_R20628
                                 R38487
                                 NM 004477
                                               Hs.203772
                                                            FSHD region gene 1
         101341 L76159
                                 AA876002
                                               Hs.120551
                                                            toli-tike receptor 10
         115455 RC_AA285068
                                                            Homo saplens clone 24775 mRNA sequence
         111192 RC_AA477748
                                 AW021968
                                               Hs.109438
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         129385 RC_AA235604
                                 AA172106
                                               Hs.110950
                                                            Rag C protein
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122105 RC_AA432278
                                 AW970209
                                               Hs.111805
                                                            ESTs
                                 AW241685
                                               Hs.98699
                                                            ESTs
         121324 RC_AA404229
120938 RC_AA386260
                                 AA404229
                                               Hs.97842
                                                            EST
                                 AA386260
                                               Hs.104632
40
                                                            gb:zs10a06.s1 NCI_CGAP_GCB1 Homo sapiens cDNA clone IMAGE:684754 3', mRNA
         115001 RC_AA251376
                                 AA251376
         sequence.
         124799 RC_R45088
                                                             gb:yg38g04.s1 Soares infant brain 1NIB Homo sapiens cDNA clone IMAGE:34896 3', mRNA
                                 R45088
         sequence.
                                                             ESTs
         122724 RC_AA457395
                                 AA457395
                                               Hs 99457
45
         117791 RC_N48325
                                 N48325
                                               Hs.93956
                                                             EST
                                                             gb:zw33a02.s1 Soares ovary tumor NbHOT Homo saptens cDNA clone IMAGE:771050 3*
         121895 RC_AA427396
                                 AA427396
         similar to contains Alu repetitive element contains MER12
                                                            t2 MER12 repetitive element; mRNA sequence.
                                                            gb:zm05c09.s1 Stratagene comeal stroma (937222) Homo sapiens cDNA clone IMAGE:513232
         108244 RC_AA062839
                                 AA062839
         3°, mRNA sequence.
         117852 RC_N49408
109298 RC_AA205432
50
                                                             KIAA0853 protein
                                 AW877787
                                               Hs.136102
                                 R77854
                                               Hs.250693
                                                             Krueppel-related zinc finger protein
                                                             ESTs, Weakly similar to B34087 hypothetical protein [H.saplens]
         122432 RC_AA447400
                                 AA447400
                                               Hs.187684
                                                             gb:za55c03.s1 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:296452 3*
         124627 RC_N74625
                                 N74625
        similar to gb:M14338 VITAMIN K-DEPENDENT PROTEIN S PRECURSOR (HUMAN); contains OFR.b3 OFR repetitive element;, mRNA sequence.

115141 RC_AA258071 AA465131 Hs.64001 Homo saplens clone 25218 mRNA sequence
55
         128636 U49065
                                 U49065
                                               Hs.102865
                                                            interleukin 1 receptor-like 2
         115373 RC_AA282197
                                 AA664862
                                               Hs.181022
                                                             CGI-07 protein
         114651 RC_AA101400
                                 AA101400
                                               Hs.189960
                                                             ESTs
         132796 RC_AA180487
                                 NM_006283
                                               Hs.173159
                                                             transforming, acidic colled-coll containing protein 1
60
         103749 RC_N35583
                                 AL135301
                                               Hs.8768
                                                             hypothetical protein FLJ10849
         107328 T83444
                                 AW959891
                                               Hs.76591
                                                             KIAA0887 protein
                                                             DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 16
         115349 RC_AA281563
                                 AF121176
                                               Hs.12797
                                                             gb:yf11e09.s1 Soares fetal liver spleen 1NFLS Homo saplens cDNA clone IMAGE:126568 3'
         111490 RC_R06862
                                 R06862
         similar to contains L1 repetitive element; mRNA sequence
65.
         103763 AA085354
                                 AA085291
                                                             gb:zn01g06.s1 Stratagene colon HT29 (937221) Homo sapiens cDNA clone 3' similar to
         contains Alu repetitive element, mRNA sequence
         118791 RC_N75520
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                                                             ESTs, Moderately similar to B34087 hypothetical protein [H.sapiens]
                                 N75520
         116644 RC_F03032
                                                             ESTs, Wealty similar to B34087 hypothetical protein [H.sapiens]
                                  F03032
                                               Hs.290278
                                                             ESTs, Highly similar to CSA_HUMAN COCKAYNE SYNDROME WD-REPEAT PROTEIN CSA
         116823 RC_H56485
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                                               Hs.143542
70
         [H.saptens]
         108940 RC_AA148603
                                                             gb:zo09e04.s1 Stratagene neuroepithelium NT2RAMI 937234 Homo sapiens cDNA clone
                                 AA148603
         IMAGE:567198 3", mRNA sequence.
         112218 RC_R50057
116557 RC_D20572_I
                                  R50057
                                               Hs.272251
                                                             Homo sapiens mRNA; cDNA DKFZp586M1418 (from clone DKFZp586M1418)
                                  D20572
                                               Hs.90171
75
         133649 U25849
                                  U25849
                                               Hs.75393
                                                             acid phosphatase 1, soluble
                                                             ESTs, Moderately similar to A45010 X-linked retinopathy protein [H.sapiens]
         131745 RC_C20746
                                  AJ828559
                                               Hs.31447
```

		RC_H43879	H43879		gb:yo69h09.s1 Soares breast 3NbHBst Homo sapiens cDNA clone IMAGE:183233 3*, mRNA
	sequenc		A 4054540	11- 07000	POT
		RC_AA251548	AA251548	Hs.87886	EST
5		RC_AA598500	H29882	Hs.162614	ESTs
,		RC_AA347919	AA347919	Hs.96889	EST
		AA018298	AA018298	Hs.103332	ESTs
		RC_AA412491	AF025771	Hs.50123	zinc finger protein 189
		RC_N67946	N67946	Hs.117569	ESTs
10		RC_AA058946	AB020700	Hs.3830	KIAA0893 protein
10		RC_AA194568_i		Hs.85938	EST
		RC_AA252794	AA252794	Hs.88009	ESTs
	420208		BE276055	Hs.95972	silver (mouse homolog) like
		AC002045_xpt1	R66740	Hs.110613	KIAA0220 protein
15	128546		NM_003478	Hs.101299	culin 5
13		RC_R40697	R40697	Hs.76666	C9orf10 protein
		RC_R73588	R73588	Hs.101533	ESTs
		AA410345	AF255910	Hs.54650	junctional adhesion molecule 2
		AA018418	AW137912	Hs.227583	Homo saplens chromosome X map Xp11.23 L-type calcium channel alpha-1 subunit
20					nplate sequence; and JM1 protein, JM2 protein, and Hb2E genes, complete cds
20		AA089688	BE244667 W90022	Hs.296155	CGI-100 protein
		W90022	VV9UUZZ	Hs.186809	ESTs, Highly similar to LCT2_HUMAN LEUKOCYTE CELL-DERIVED CHEMOTAXIN 2
		RSOR [H.saplens]	AMPOODAA	U- 002507	TOT-
		AA435512	AW298244 BE258532	Hs.293507	ESTs CTR continue
25		RC_AA404494		Hs.251871	CTP synthase
23		RC_AA278529_I		Hs.172052	serine/threonine kinase 18
		RC_AA342828_s		Hs.73734	glycoprotein V (platelet)
		RC_AA044644	AW247017	Hs.36978	melanoma antigen, family A, 3
		RC_AA044644	T07568	Hs.137158	ESTs .
30		RC_AA196729_i		Hs.190213	ESTS
20		RC_AA196729_i RC_AA025858		Hs.334825	Homo saplens cDNA FLJ14752 fis, clone NT2RP3003071
		RC_AA025858	AW247017 N75346	Hs.36978 Hs.82906	melanoma antigen, family A, 3 CDC20 (cell division cycle 20, S. cerevisiae, homolog)
			Al949109	Hs.246885	hypothetical protein FLI20783
		RC_N52271	D31139	Hs.154103	LIM protein (similar to rat protein kinase C-binding enigma)
35		RC_N68399	NM_003528	Hs.2178	H2B histone family, member Q
55		RC_AA098874	AB037850	Hs.16621	DKFZP434I116 protein
		RC_AA279667_s		Hs.297939	cathepsin B
		RC_H22556	W27893	Hs.150580	putative translation initiation factor
		RC_N45979_s	BE296690	Hs.288173	Homo sapiens cDNA: FLJ21747 fis, clone COLF5160, highly similar to AF182198 Homo sapiens
40		in 2 long isoform (f		113.200173	Truino sapiens correct caz 17-17 hs, cubite occid 5 foot, highly shindar to Al 102 foot for his separats
10		RC_AA431288_s		Hs.95327	CD3D antigen, delta polypeptide (TiT3 complex)
		RC_AA609862	BE280456	Hs.80248	RNA-binding protein gene with multiple splicing
		RC_N35583	AW994032	Hs.8768	hypothetical protein FLJ10849
	124541		AF112222	Hs.44499	pinin, desmosome associated protein
45		AA203147	AL020996	Hs.8518	selenoprotein N
		RC_W93092	AW175787	Hs.334841	selenium binding protein 1
		RC_W93092	AW516428	Hs.78687	neutral sphingomyelinase (N-SMase) activation associated factor
		RC_N58561_s	L04288	Hs.297939	cathepsin B
		RC_W93092	AW516428	Hs.78687	neutral sphingomyelinase (N-SMase) activation associated factor
50		RC_W69385_s	BE543412	Hs.250505	retinolc acid receptor, alpha
,,,		RC_R22947	R23053	NA	Hu01 Chip Redos
		RC_N38959_f	AL042444	Hs.62402	p21/Cdo42/Rac1-activated kinase 1 (yeast Ste20-related)
		RC_N38959_f	A1678638	Hs.6456	chaperonin containing TCP1, subunit 2 (beta)
		RC_H73050_s	AA744902	Hs.107767	hypothetical protein PRO1489
55			X54534	Hs.278994	Rhesus blood group, CoEe antigens
					A 141A

TABLE 1A

Table 1A shows the accession numbers for those pkeys lacking unigenelD's for Tables 1. The pkeys in Table 7 lacking unigenelD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligonucteotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Aligument Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" only many continuous conti

10 Pkey: Unique Eos probeset identifier number CAT number: Gene cluster number

Accession: Genbank accession numbers

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15
        Pkey CAT Number Accession
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                            AA083256
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                            AA100796 AF020589 AA074629 AA075946 AA100849 AA085347 AA126309 AA079311 AA079323 AA085274
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                            T69981 T69924 AA078476
        125066 1814993 1
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        116661 1532859_1
                            R61504 F04247
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        125104 413347_1
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                            AA098878 W88942
        116845 393481_1
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                            AI652535 BE465762 AA206331 AW451866 AA471088 AA206342 AA204834 AA206100 AW021661 AA332922 N66048
                            AA703396 H92278 AW139734 H92683 U87589 U87595 H69001 U87594 BE466420 Al624817 BE466611 Al206344 AA574397
35
                            AA348354 AJ493192
        118584 532052_1
                            AW136928 Al685655 BE218584 BE465078 N68963 AA975338 BE147199 N76377
                            AA075998 AA075999 AA070986 AA070896 AA129207 AA078942 AA070783 AA078941
        103743 112194_1
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                            AA079267 AA076003
        103746 113452 1
                            AA075000 AA081876
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        103761 114208 1
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        103763 48290 6
                            AA085291 AA085354
                            F02951 Z40892 F04711
        120209 1531817_1
                            AA179656 AA182626 AA182603
        120284 158963_1
        112540 1605263_1
                            R69751 R70467 H69771 H80879 H80878
45
                            Z41572 R39330
        111904 1719336_1
        121059 273450 1
                            AA393283 AA398628
        121094 275729_1
                            AA402505 AA398900
                            AW602528 BE073859 Z38412
        114105 1182096_1
        130091 23961_-3
                            W88999
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               296527_1
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        122264
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                            R23053 R79884 R76271
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        130529
        108309 111495_1
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                                       AA128980
        125017 genbank_T68875
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60
        125063 genbank_T85352
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        125064 genbank_T85373
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        125118 149288_1
                            R10606 T97620 AA576309
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        102269
        125150 NOT_FOUND_entrez_W38240
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                                                 W38240
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                                       H43879
        118111 genbank_N55493
                                       N55493
        118129 genbank_N57493
                                       N57493
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TABLE 2:

Unique Eos probeset identifier number Accession number used for previous patent filings Exemplar Accession number, Genbank accession number Unigene number Unigene gene title Pkey: Accession: ExAccn; UnigeneID: Unigene Title; 5

	Ougene	Time, O	ingene gene uu	6	
10	Pkey	Accession	ExAccn	UnigenelD	UnigeneTitle
	100420	100420	D86983	Hs.118893	Melanoma associated gene
	100484	100484	NM_005402	Hs.288757	v-ral simian leukemla viral oncogene hom
	100991	100991	J03836	Hs.82085	serine (or cysteine) proteinase inhibito
15		101168	NM_005308	Hs.211569	G protein-coupled receptor kinase 5
		101261	D30857	Hs.82353	protein C receptor, endothelial (EPCR)
		101447	M21305		gb:Human alpha satellite and satellite 3
		101543	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
20		101560	AW958272	Hs.347326	intercellular adhesion molecule 2
20		101714 101838	M68874 BE243845	Hs.211587 Hs.75511	phospholipase A2, group IVA (cytosolic, connective tissue growth factor
		102012	BE259035	Hs.118400	singed (Drosophila)-like (sea urchin fas
		102012	NM_000107	Hs.77602	damage-specific DNA binding protein 2 (4
		102283	AW161552	Hs.83381	quanine nucleotide binding protein 11
25		102564	U59423	Hs.79067	MAD (mothers against decapentaplegic, Dr
		102759	NM_005100	Hs.788	A kinase (PRKA) anchor protein (gravin)
		102804	NM_002318	Hs.83354	lysyl oxidase-like 2
	102898	102898	NM_002205	Hs.149609	integrin, alpha 5 (fibronectin receptor,
		103036	M13509	Hs.83169	matrix metalloproteinase 1 (Interstitial
30		103095	NM_005424	Hs.78824	tyrosine kinase with immunoglobulin and
		103166	AA159248	Hs.180909	peroxiredoxin 1
		103280	U84722	Hs.76206	cadherin 5, type 2, VE-cadherin (vascula
		103850	AA187101	Hs.213194	hypothetical protein MGC10895
35		104592	AW630488	Hs.25338 Hs.10031	protease, serine, 23 · KIAA0955 protein
33		104786 104865	AA027167 T79340	Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
		104952	AW076098	Hs.345588	desmoplakin (DPI, DPII)
		105178	AA313825	Hs.21941	AD036 protein
		105330	AW338625	Hs.22120	ESTs
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		105977	AK001972	Hs.30822	hypothetical protein FLJ11110
		106031	X64116	Hs.171844	Homo saptens cDNA: FLJ22296 fis, clone H
		106155	AA425414	Hs.33287	nuclear factor I/B
AE		106423	AB020722	Hs.16714	Rho guanine exchange factor (GEF) 15
45		107174	BE122762	Hs.25338	ESTS
		107295 108756	AA186629 AA127221	Hs.80120 Hs.117037	UDP-N-acetyl-alpha-D-galactosamine:polyp ESTs
		108888	AA135606	Hs.189384	gb:zl10a05.s1 Soares_pregnant_uterus_NbH
		109166	AA219691	Hs.73625	RAB6 interacting, kinesin-like (rabkines
50		109768	F06838	Hs.14763	ESTs
Ţ		110906	AA035211	Hs.17404	ESTs
		111006	BE387014	Hs.166146	Homer, neuronal immediate early gene, 3
	111133	111133	AW580939	Hs.97199	complement component C1q receptor
		113073	N39342	Hs.103042	microtubule-associated protein 1B
55		113923	AW953484	Hs.3849	hypothetical protein FLJ22041 similar to
		115061	AI751438	Hs.41271	Homo sapiens mRNA full length insert cDN
		115145	AA740907	Hs.88297 Hs.94761	ESTs
		115947 116339	R47479 AK000290	Hs.44033	KIAA1691 protein dipentidyl peptidase 8
60		116589	Al557212	Hs.17132	ESTs, Moderately similar to I54374 gene
00		117023	AW070211	Hs.102415	Homo sapiens mRNA; cDNA DKFZp586N0121 (f
	117563	117563	AF055634	Hs.44553	unc5 (C.elegans homolog) c
	118475	118475	N66845	,,,,,,,,,,	gb:za46c11.s1 Soares fetal liver spleen
	119073	119073	BE245360	Hs.279477	ESTs
65	119174	119174	R71234		gb:yi54c08.s1 Soares placenta Nb2HP Homo
	119416	119416	T97186		gb:ye50h09.s1 Soares fetal liver spleen
	121335	121335	AA404418		gb:zw37e02.s1 Soares_total_fetus_Nb2HF8_
	123160	123160	AA488687	Hs.284235	ESTs, Weakly similar to 138022 hypotheti
70	123523	123523	AA608588		gb:ae54e06.s1 Stratagene lung carcinoma
70	123964	123964	C13961	U. 000777	gb:C13961 Clontech human aorta polyA+ mR
	124315	124315 124669	NM_005402 Al571594	Hs.288757 Hs.102943	v-ral simian leukemia viral oncogene hom hypothetical protein MGC12916
	124009	124875	AIS7 1594 AI887664	Hs.285814	sprouty (Drosophila) homolog 4
	124010	125103	AA570056	Hs.122730	ESTs. Moderately similar to KIAA1215 pro
75	125565	125565	R20840	, N. 1221 00	gb:yg05c08.r1 Scares infant brain 1NIB H
	12,000				Gardon and an

	126511 126511	T92143	Hs.57958	ECE TAT Introdulin related protein
	126649 126649	AA001860	Hs.279531	EGF-TM7-latrophillin-related protein ESTs
	449602 449602	AA001860	Hs.279531	ESTs
	127402 127402	AA358869	Hs.227949	SEC13 (S. cerevisiae)-like 1
5	128992 128992	H04150	Hs.107708	ESTs .
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	129371 129371	X06828	Hs.110802	von Willebrand factor
	129765 129765	M86933	Hs.1238	amelogenin (Y chromosome)
	129884 129884	AF055581	Hs.13131	lysosomal
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	130828 130828	AW631469	Hs.203213	ESTs
	131080 131080	NM_001955	Hs.2271	endothelin 1
	131182 131182	A1824144	Hs.23912	ESTs ·
1.5	131573 131573	AA040311	Hs.28959	ESTs
15	131756 131756	AA443966	Hs.31595	ESTs
	131881 131881	AW361018	Hs.3383	upstream regulatory element binding prot
	132083 132083	BE386490	Hs.279663	Pirtn .
	132358 132358	NM_003542	Hs.46423	H4 histone family, member G
20	132456 132456 132676 132676	AB011084 N92589	Hs.48924 Hs.261038	KIAA0512 gene product; ALEX2 ESTs, Wealdy similar to I38022 hypotheti
20	132718 132718	NM_004600	Hs.554	Sjogren syndrome antigen A2 (60kD, ribon
	132760 132760	AA125985	Hs.56145	thymosin, beta, identified in neuroblast
	132968 132968	AF234532	Hs.61638	myosin X
	133061 133061	Al186431	Hs.296638	prostate differentiation factor
25	133161 133161	AW021103	Hs.6631	hypothetical protein FLJ20373
	133260 133260	AA403045	Hs.6906	Homo sapiens cDNA: FLJ23197 fis, clone R
	133491 133491	BE619053	Hs.170001	eukaryotic translation initiation factor
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	133614 133614	NM_003003	Hs.75232	SEC14 (S. cerevisiae)-like 1
30	133691 133691	M85289	Hs.211573	heparan suifate proteoglycan 2 (periecan
	133913 133913	AU076964	Hs.7753	calumenin
	133985 133985	L34657	Hs.78146	platelet/endothelial cell adhesion molec
	134088 134088	Al379954	Hs.79025	KIAA0096 protein
35	134299 134299	AW580939	Hs.97199	complement component C1q receptor
33	116470 116470	Al272141 AW968058	Hs.83484 Hs.92381	SRY (sex determining region Y)-box 4 nudix (nucleoside diphosphate linked moi
	134989 134989 135073 135073	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
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	100405 100405	AW291587	Hs.82733	nidogen 2
	100455 100455	AW888941	Hs.75789	N-myc downstream regulated
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	100658 100658	U56725	Hs.180414	heat shock 70kD protein 2
45	100718 100718	BE295928	Hs.75424	inhibitor of DNA binding 1, dominant neg
	100828 100828	AL048753	Hs.303649	small inducible cytokine A2 (monocyte ch
	100991 100991	J03836	Hs.82085	serine (or cystelne) proteinase inhibito
	101110 101110	Al439011	Hs.86386	myeloid cell teukemla sequence 1 (BCL2-r
50	101156 101156	AA340987	Hs.75693	prolylcarboxypeptidase (angiotensinase C
30	101184 101184	NM_001674	Hs.460	activating transcription factor 3
	101317 101317 101345 101345	L42176	Hs.8302 Hs.152175	four and a half LIM domains 2 calcitonin receptor-like
	101345 101345	NM_005795 BE410405	Hs.76288	calpain 2, (m/ll) large subunit
	101496 101496	X12784	Hs.119129	collagen, type IV, alpha 1
55	101543 101543	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
55	101560 101560	AW958272	Hs.347326	intercellular adhesion molecule 2
	101592 101592	AF064853	Hs.91299	guanine nucleotide binding protein (G pr
	101634 101634	AV650262	Hs.75765	GRO2 oncogene
	101682 101682	AF043045	Hs.81008	filamin B, beta (actin-binding protein-2
60	101720 101720	M69043	Hs.81328	nuclear factor of kappa light polypeptid
	101744 101744	Al879352	Hs.118625	hexokinase 1
	101837 101837	M92843	Hs.343586	zinc finger protein homologous to Zfp-36
	101840 101840	AA236291	Hs.183583	serine (or cysteine) proteinase inhibito
	101864 101864	BE392588	Hs.75777	transgelin
65	101966 101966	X96438	Hs.76095	immediate early response 3
	102013 102013	BE616287	Hs.178452	catenin (cadherin-associated protein), a
	102059 102059	A1752666	Hs.76669	nicotinamide N-methyltransferase
	102283 102283	AW161552	Hs.83381	guanine nucleotide binding protein 11
70	102378 102378 102460 102460	AU076887 U48959	Hs.28491 Hs.211582	spermidine/spermine N1-acetyltransferase myosin, light polypeptide kinase
, 0	102499 102499	BE243877	Hs.76941	ATPase, Na+/K+ transporting, beta 3 poly
	102499 102499	R97457	Hs.63984	cadherin 13, H-cadherin (heart)
	102589 102589	AU076728	Hs.8867	cysteine-rich, angiogenic inducer, 61
	102645 102645	AL119566	Hs.6721	lysosomal
75	102693 102693	AA532780	Hs.183684	eukaryotic translation initiation factor
	102759 102759	NM_005100	Hs.788	A kinase (PRKA) anchor protein (gravin)

	102882 102882	Al767736	Hs.290070	gelsolin (amyloldosis, Finnish type)
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	102960 102960	Al904738	Hs.76053	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep
_	103020 103020	X53416	Hs.195464	filamin A, alpha (actin-binding protein-
5	103036 103036	M13509	Hs.83169	matrix metalloproteinase 1 (interstitial
	103080 103080 103138 103138	AU077231 X65965	Hs.82932	cyclin D1 (PRAD1: parathyroid adenomatos gb:H.sapiens SOD-2 gene for manganese su
	103195 103195	AA351647	Hs.2642	eukaryotic translation elongation factor
	103371 103371	X91247	Hs.13046	thioredoxin reductase 1
10	103471 103471	Y00815	Hs.75216	protein tyrosine phosphatase, receptor t
	104447 104447	AW204145	Hs.156044	ESTs
	104783 104783 104865 104865	AA533513 T79340	Hs.93659 Hs.22575	protein disulfide isomerase related prot B-cell CLL/lymphoma 6, member B (zinc fi
	104894 104894	AF065214	Hs.18858	phospholipase A2, group IVC (cytosolic,
15	105113 105113	AB037816	Hs.8982	Homo sapiens, done IMAGE:3506202, mRNA,
	105196 105196	W84893	Hs.9305	anglotensin receptor-like 1
	105263 105263	AW388633	Hs.6682	solute carrier family 7, (cationic amino
	105330 105330 105492 105492	AW338625 Al805717	Hs.22120 Hs.289112	ESTs CGI-43 protein
20	105594 105594	AB024334	Hs.25001	tyrosine 3-monooxygenase/tryptophan 5-mo
	105732 105732	AW504170	Hs.274344	hypothetical protein MGC12942
	105882 105882	W46802	Hs.81988	disabled (Drosophila) homolog 2 (mitogen
	106031 106031 106222 106222	X64116	Hs.171844	Homo sapiens cDNA: FLJ22296 fis, clone H Homo sapiens clone FLB9213 PRO2474 mRNA,
25	106263 106263	AA356392 W21493	Hs.21321 Hs.28329	hypothetical protein FLJ14005
23	106366 106366	AA186715	Hs.336429	RIKEN cDNA 9130422N19 gene
	106634 106634	W25491	Hs.288909	hypothetical protein FLJ22471
	106793 106793	H94997	Hs.16450	ESTs
30	106842 106842 106890 106890	AF124251 AA489245	Hs.26054 Hs.88500	novel SH2-containing protein 3 mitogen-activated protein kinase 8 Inter
50	106974 106974	AJ817130	Hs.9195	Homo sapiens cDNA FLJ13698 fis, clone PL
•	107061 107061	BE147611	Hs.6354	stromal cell derived factor receptor 1
	107216 107216	D51069	Hs.211579	melanoma cell adhesion molecule
35	107444 107444	W28391	Hs.343258	proliferation-associated 2G4, 38kD ESTs
33	108507 108507 108931 108931	Al554545 AA147186	Hs.68301	gb:zo38d01.s1 Stratagene endothetial cel
	109195 109195	AF047033	Hs.132904	solute carrier family 4, sodium bicarbon
	109456 109456	AW956580	Hs.42699	ESTs
40	110411 110411	AW001579	Hs.9645	Homo sapiens mRNA for KIAA1741 protein,
40	110906 110906 111091 111091	AA035211 AA300067	Hs.17404 Hs.33032	ESTs hypothetical protein DKFZp434N185
	111378 111378	AW160993	Hs.326292	hypothetical gene DKFZp434A1114
	111769 111769	AW629414	Hs.24230	ESTs
4.5	112951 112951	AA307634	Hs.6650	vacuolar protein sorting 45B (yeast homo
45	113195 113195 113542 113542	H83265 H43374	Hs.8881 Hs.7890	ESTs, Weakly similar to S41044 chromosom Homo sapians mRNA for KIAA1671 protein,
	113847 113847	NM_005032	Hs.4114	plastin 3 (T isoform)
	113947 113947	W84768		gb:zh53d03.s1 Soares_fetal_liver_spleen_
50	115061 115061	Al751438	Hs.41271	Homo sapiens mRNA full length insert cDN
50	115870 115870	NM_005985	Hs.48029	snail 1 (drosophila homolog), zinc finge ESTs
	116228 116228 116314 116314	A1767947 A1799104	Hs.50841 Hs.178705	Homo sapiens cDNA FLJ11333 fis, clone PL
	117023 117023	AW070211	Hs.102415	Homo saplens mRNA; cDNA DKFZp586N0121 (f
	117156 117156	W73853		ESTs
55	117280 117280	M18217	Hs.172129	Homo sapiens cDNA: FLJ21409 fis, clone C
	119866 119866 121314 121314	AA496205 W07343	Hs.193700 Hs.182538	Homo saplans mRNA; cDNA DKFZp58610324 (f phospholipid scramblase 4
	121822 121822	AI743860	110.102000	metallothionein 1E (functional)
	122331 122331	AL133437	Hs.110771	Homo saplens cDNA: FLJ21904 fis, clone H
60	123160 123160	AA488687	Hs.284235	ESTs, Wealdy similar to 138022 hypotheti
	124059 124059 124358 124358	BE387335 AW070211	Hs.283713 Hs.102415	ESTs, Weakly similar to S64054 hypotheti Homo sapiens mRNA; cDNA DKFZo586N0121 (f
	124726 124726	NM_003654	Hs.104576	carbohydrate (keratan sulfate Gal-6) sul
	125167 125167	AL137540	Hs.102541	netrin 4
65	125307 125307	AW580945	Hs.330466	ESTs
	107985 107985	T40064	Hs.71968	Homo sapiens mRNA; cDNA DKFZp564F053 (fr Homo sapiens mRNA; cDNA DKFZp564F053 (fr
	125598 125598 413731 413731	T40064 BE243845	Hs.71968 Hs.75511	connective tissue growth factor
	116024 116024	AA088767	Hs.83883	transmembrane, prostate androgen induced
70	418000 418000	AA932794	Hs.83147	guanine nucleotide binding protein-like
	126399 126399	AA088767	Hs.83883	transmembrane, prostate androgen induced
	127566 127566 128453 128453	AI051390 X02761	Hs.116731 Hs.287820	ESTs fibranectin 1
	128515 128515	BE395085	Hs.10086	type I transmembrane protein Fn14
75	128623 128623	BE076608	Hs.105509	CTL2 gene
	128669 128669	W28493	Hs.180414	heat shock 70kD protein 8

	420044 420044	A18/007/04	Un 40740E	alasmalasmas varieta accepiated amtain
	128914 128914 129188 129188	AW867491 NM_001078	Hs.107125 Hs.109225	plasmalemma vesicle associated protein vascular cell adhesion molecule 1
	129265 129265	AA530892	Hs.171695	dual specificity phosphatase 1
	129468 129468	AW410538	Hs.111779	secreted protein, acidic, cysteline-rich
5	101838 101838	BE243845	Hs.75511	connective tissue growth factor
•	129619 129619	AA209534	Hs.284243	tetraspan NET-6 protein
	129762 129762	AA453694	Hs.12372	tripartite motif protein TRIM2
	130018 130018	AA353093		metallothionein 1L
	130178 130178	U20982	Hs.1516	Insulin-like growth factor-binding prote
10	130431 130431	AW505214	Hs.155560	calnexin
	130553 130553	AF062649	Hs.252587	pituitary tumor-transforming 1
	130639 130639	Al557212	Hs.17132	ESTs, Moderately similar to I54374 gene
	130686 130686	BE548267	Hs.337986	Homo saplens cDNA FLJ10934 fis, clone OV
	130818 130818	AW190920	Hs.19928	hypothetical protein SP329
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	131080 131080	NM_001955	Hs.2271	endothelin 1
	131091 131091	AJ271216	Hs.22880	dipeptidylpeptidase (!)
	131182 131182	AI824144	Hs.23912	ESTs
20	131319 131319	NM_003155	Hs.25590	stanniocaldin 1
20	131328 131328	AW939251	Hs.25647	v-fos FBJ murine osteosarcoma viral onco v-fos FBJ murine osteosarcoma viral onco
	131328 131328 131555 131555	AW939251 T47364	Hs.25647 Hs.278613	interferon, alpha-inducible protein 27
	131573 131573	AA040311	Hs.28959	ESTs
	131756 131756	AA443966	Hs.31595	ESTs
25	131909 131909	NM_016558	Hs.274411	SCAN domain-containing 1
	132046 132046	Al359214	Hs.179260	chromosome 14 open reading frame 4
	132151 132151	BE379499	Hs.173705	Homo sapiens cDNA: FLJ22050 fis, clone H
	132187 132187	AA235709	Hs.4193	DKFZP586O1624 protein
	132314 132314	AF112222	Hs.323806	pinin, desmosome associated protein
30	132398 132398	AA876616	Hs.16979	ESTs, Weakly similar to A43932 mucin 2 p
	132490 132490	NM_001290	Hs.4980	LIM domain binding 2
	132546 132546	M24283	Hs.168383	intercellular adhesion molecule 1 (CD54)
	132716 132716	BE379595	Hs.283738	casein kinase 1, alpha 1
25	132883 132883	AA373314	Hs.5897	Homo sapiens mRNA; cDNA DKFZp586P1622 (f
35	132989 132989	AA480074	Hs.331328	hypothetical protein FLJ13213
	133071 133071	BE384932	Hs.64313	ESTs, Weakly similar to AF257182 1 G-pro
	133099 133099	W16518	Hs.279518	amyloid beta (A4) precursor-like protein
	133149 133149	AA370045 AB037715	Hs.6607 Hs.183639	AXIN1 up-regulated hypothetical protein FLJ10210
40	133200 133200 133260 133260	AA403045	Hs.6906	Homo sapiens cDNA: FLJ23197 fls, clone R
TU	133349 133349	AW631255	Hs.8110	L-3-hydroxyacyl-Coenzyme A dehydrogenase
	133398 133398	NM_000499	Hs.72912	cytochrome P450, subfamily I (aromatic c
	133454 133454	BE547647	Hs.177781	hypothetical protein MGC5618
	133491 133491	BE619053	Hs.170001	eukaryotic translation initiation factor
45	133517 133517	NM_000165	Hs.74471	gap junction protein, alpha 1, 43kD (con
	133538 133538	NM_003257	Hs.74614	tight junction protein 1 (zona occludens
	133584 133584	D90209	Hs.181243	activating transcription factor 4 (tax-r
	133617 133617	BE244334	Hs.75249	ADP-ribosylation factor-like 6 Interacti
~~	133671 133671	AW503116	Hs.301819	zinc finger protein 146
50	133681 133681	A1352558		tyrosine 3-monooxygenase/tryptophan 5-mo
	133730 133730	BE242779	Hs.179526	upregulated by 1,25-dihydroxyvitamin D-3
	133802 133802	AW239400	Hs.76297	G protein-coupled receptor kinase 6
	133838 133838	BE222494	Hs.180919	inhibitor of DNA binding 2, dominant neg
55	133889 133889 133975 133975	U48959 C18356	Hs.211582 Hs.295944	rnyosin, light polypeptide kinase tissue factor pathway inhibitor 2
55	134039 134039	NM_002290	Hs.78672	laminin, alpha 4
	134081 134081	AL034349	Hs.79005	protein tyrosine phosphatase, receptor t
	134203 134203	AA161219	Hs.799	diphtheria toxin receptor (heparin-bindi
	134299 134299	AW580939	Hs.97199	complement component C1g receptor
60	134339 134339	R70429	Hs.81988	disabled (Drosophila) homolog 2 (mitogen
••	134381 134381	AI557280	Hs.184270	capping protein (actin filament) muscle
	134416 134416	X68264	Hs.211579	melanoma cell adhesion molecule
	134558 134558	NM_001773	Hs.85289	CD34 antigen
	134983 134983	D28235	Hs.196384	prostaglandin-endoperoxide synthase 2 (p
65	135052 135052	AL136653	Hs.93675	decidual protein induced by progesterone
	135069 135069	AA876372	Hs.93961	Homo saplens mRNA; cDNA DKFZp667D095 (fr
	135073 135073	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
	135196 135196	C03577	Hs.9615	myosin regulatory light chain 2, smooth
70	134404 134404	AB000450	Hs.82771	vaccinia related kinase 2
70	100082 100082	AA130080	Hs.4295	proteasome (prosome, macropain) 26S subu
	130150 130150	BE094848 AB011169	Hs.15113	homogentisate 1,2-dioxygenase (homogenti
	130839 130839 100113 100113	NM_001269	Hs.20141	similar to S. cerevisiae SSM4 chromosome condensation 1
	100129 100129	AA469369	Hs.84746 Hs.5831	tissue inhibitor of metalloproteinase 1
75	100169 100169	AL037228	Hs.82043	D123 gene product
. 5	100190 100190	M91401	Hs.178658	RAD23 (S. cerevisiae) hornolog B
	.00.00			(a. anarona) intimon

	400214	400044	DOCEDO	Un 4920E0	DEAD(U (Asp. Ch. Alb. Asp./Uis) hav pohmon
	100211		D26528 NM_012288	Hs.123058 Hs.153954	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep TRAM-like protein
		100248	NM_015156	Hs.78398	KIAA0071 protein
		100262	D38500	Hs.278468	postmeiotic segregation increased 2-like
5	100281		AF091035	Hs.184627	KIAA0118 protein
3		100327	D55640	113.101021	gb:Human monocyte PABL (pseudoautosomal
		134495	D63477	Hs.84087	KIAA0143 protein
		135152	M96954	Hs.182741	TIA1 cytotoxic granule-associated RNA-bl
		100372	NM_014791	Hs.184339	KIAA0175 gene product
10		100394	D84284	Hs.66052	CD38 antigen (p45)
••		100418	D86978	Hs.84790	KIAA0225 protein
	134347		AF164142	Hs.82042	solute carrier family 23 (nucleobase tra
		100438	AA013051	Hs.91417	topoisomerase (DNA) II binding protein
	100481		X70377	Hs.121489	cystatin D
15	100591	100591	NM_004091	Hs.231444	Homo saplens, Similar to hypothetical pr
		100662	Al368680	Hs.816	SRY (sex determining region Y)-box 2
	100905	100905	L12260	Hs.172816	neuregulin 1
		100950	AF128542	Hs.166846	polymerase (DNA directed), epsilon
	135407	135407	J04029	Hs.99936	keratin 10 (epidermolytic hyperkeratosis
20	131877	131877	J04088	Hs.156346	topoisomerase (DNA) II alpha (170kD)
	134786	134786	T29618	Hs.89640	TEK tyrosine kinase, endothelial (venous
	134078	134078	L08895	Hs.78995	MADS box transcription enhancer factor 2
	134849	134849	BE409525	Hs.902	neurofibromin 2 (bilateral acoustic neur
	101152	101152	Al984625	Hs.9884	spindle pole body protein
25	131687	131687	BE297635	Hs.3069	heat shock 70kD protein 9B (mortalin-2)
	421155	421155	H87879	Hs.102267	lysyl oxidase
	133975	133975	C18356	Hs.295944	tissue factor pathway inhibitor 2
	130155	130155	AA101043	Hs.151254	kallikrein 7 (chymotryptic, stratum com
	132813	132813	BE313625	Hs.57435	solute carrier family 11 (proton-coupled
30	101300	101300	BE535511		transmembrane trafficking protein
	130344	130344	AW250122	Hs.154879	DiGeorge syndrome critical region gene D
	101381	101381	AW675039	Hs.1227	aminolevulinate, delta-, dehydratase.
		133780	AA557660	Hs.76152	decorin
25		101447	M21305		gb:Human alpha satellite and satellite 3
35		101470	NM_000546	Hs.1846	tumor protein p53 (Li-Fraumeni syndrome)
		101478	NM_002890	Hs.758	RAS p21 protein activator (GTPase activa
		133519	AW583062	Hs.74502	chymotrypsinogen B1
		134116	R84694	Hs.79194	cAMP responsive element binding protein
40		130174	M29551	Hs.151531	protein phosphatase 3 (formerly 2B), cat
40		132983	M30269	11. 0050	nidogen (enactin)
		101543	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
		101620	S55271	Hs.247930	Epsilon , IgE
		133595	AA393273	Hs.75133	transcription factor 6-like 1 (mitochond
45		101700	D90337 D28459	Hs.247916	natriuretic peptide precursor C ubiquitin-conjugating enzyme E2A (RAD6 h
43		134246		Hs.80612	sphingomyelin phosphodiesterase 1, acid
		133948	X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
		133948 133948	X59960 X59960	Hs.77813 Hs.77813	sphingomyelin phosphodiesterase 1, acid
		101812	BE439894	Hs.78991	DNA segment, numerous copies, expressed
50		133396	M96326	Hs.72885	azurocidin 1 (cationic antimicrobial pro
50		129026	AL120297	Hs.108043	Friend leukemia virus integration 1
		134831	AA853479	Hs.89890	pyruvate carboxylase
		134395	AA456539	Hs.8262	lysosomal
		101977	AF112213	Hs.184062	putative Rab5-interacting protein
55		101998	U01212	Hs.248153	olfactory marker protein
		102007	U02556	Hs.75307	t-complex-associated-testis-expressed 1-
		416658	U03272	Hs.79432	fibrillin 2 (congenital contractural ara
		135389	U05237	Hs.99872	fetal Alzheimer antigen
		130145	U34820	Hs.151051	mitogen-activated protein kinase 10
60		420269	U72937	Hs.96264	alpha thalassemia/mental retardation syn
		102123	NM_001809	Hs.1594	centromere protein A (17kD)
		102133	AU076845	Hs.155596	BCL2/adenovirus E1B 19kD-interacting pro
		102162	AA450274	Hs.1592	CDC16 (cell division cycle 16, S. cerevi
	427653	427653	AA159001	Hs.180069	nuclear respiratory factor 1
65	102200	102200	AA232362	Hs.157205	branched chain aminotransferase 1, cytos
	102214	102214	U23752	Hs.32964	SRY (sex determining region Y)-box 11
	131319	131319	NM_003155	Hs.25590	stanniocaldin 1
		132316	U28831	Hs.44566	KIAA1641 protein
		134365	AA568906	Hs.82240	syntaxin 3Å
70		102298	AA382169	Hs.54483	N-myc (and STAT) interactor
		302344	BE303044	Hs.192023	eukaryotic translation initiation factor
		102357	U39656	Hs.118825	mitogen-activated protein kinase kinase
	102394	102394	NM_003816	Hs.2442	a disintegrin and metalloproteinase doma
76		129521	AF071076	Hs.112255	nucleoporin 98kD
75		102251	NM_004398	Hs.41706	DEAD/H (Asp-Ghi-Ala-Asp/His) box polypep
	133746	133746	AW410035	Hs.75862	MAD (mothers against decapentaplegic, Dr

		10044045		
	132828 132828	AB014615	Hs.57710	fibroblast growth factor 8 (androgen-ind
	132828 132828	AB014615	Hs.57710	fibroblast growth factor 8 (androgen-Ind
	130441 130441	U63630	Hs.155637	protein kinase, DNA-activated, catalytic
_	129350 129350	U50535	Hs.110630	Human BRCA2 region, mRNA sequence CG006
5	130457 130457	AB014595	Hs.155976	cullin 4B
	102560 102560	R97457	Hs.63984	cadherin 13, H-cadherin (heart)
	134305 134305	U61397	Hs.81424	ubiquitin-like 1 (sentrin)
	132736 132736	AW081883	Hs.211578	Homo sapiens cDNA: FLJ23037 fis, clone L
	102663 102663	NM_002270	Hs.168075	karyopherin (importin) beta 2
10	102735 102735	AF111106	Hs.3382	protein phosphatase 4, regulatory subuni
	101175 101175	U82671	Hs.36980	melanoma antigen, family A, 2
	132164 132164	AI752235	Hs.41270	procollagen-lysine, 2-oxoglutarate 5-dio
	102826 102826	NM_007274	Hs.8679	cytosofic acyl coenzyme A thioester hydr
	102846 102846	BE264974	Hs.6566	thyroid hormone receptor interactor 13
15	134161 134161	AA634543	Hs.79440	IGF-II mRNA-binding protein 3
	302363 302363	AW163799	Hs.198365	2,3-bisphosphoglycerate mutase
	125701 125701	T72104	Hs.93194	apolipoprotein A-I
	134656 134656	Al750878	Hs.87409	thrombospondin 1
	102968 102968	AU076611	Hs.154672	methylene tetrahydrofolate dehydrogenase
20	134037 134037	A1808780	Hs.227730	integrin, alpha 6
	103023 103023	AW500470	Hs.117950	multifunctional polypeptide similar to S
	130282 130282	BE245380	Hs.153952	5' nucleotidase (CD73)
	128568 128568	H12912	Hs.274691	adenylate kinase 3
	103093 103093	S79876	Hs.44926	dipeptidytpeptidase IV (CD26, adenosine
25	129063 129063	X63094	Hs.283822	Rhesus blood group, D antigen
	133227 133227	AW977263	Hs.68257	general transcription factor IIF, polype
	103184 103184	U43143	Hs.74049	fms-related tyrosine kinase 4
	103208 103208	AW411340	Hs.31314	retinoblastoma-binding protein 7
	131486 131486	F06972	Hs.27372	BMX non-receptor tyrosine kinase
30	103334 103334	NM_001260	Hs.25283	cyclin-dependent kinase 8
	135094 135094	NM_003304	Hs.250687	transient receptor potential channel 1
	103352 103352	H09366	Hs.78853	uradi-DNA giyoosylase
	132173 132173	X89426	Hs.41716	endothelial cell-specific molecule 1
	131584 131584	AA598509	Hs.29117	purine-rich element binding protein A
35	103378 103378	AL119690	Hs.153618	HCGVIII-1 protein
	103410 103410	AA158294	Hs.295362	DR1-associated protein 1 (negative cofac
	103438 103438	AW175781	Hs.152720	M-phase phosphoprotein 6
	103452 103452	NM_006936	Hs.85119	SMT3 (suppressor of mif two 3, yeast) ho
	135185 135185	AW404908	Hs.96038	Ric (Drosophila)-like, expressed in many
40	134662 134662	NM_007048	Hs.284283	butyrophilin, subfamily 3, member A1
	103500 103500	AW408009	Hs.22580	alkylgiycerone phosphate synthase
	132084 132084	NM_002267	Hs.3886	karyopherin alpha 3 (importin alpha 4)
	133152 133152	Z11695	Hs.324473	mitogen-activated protein kinase 1
	103612 103612	BE336654	Hs.70937	H3 histone family, member A
45	103692 103692	AW137912	Hs.227583	Homo sapiens chromosome X map Xp11.23 L-
	129796 129796	BE218319	Hs.5807	GTPase Rab14
	132683 132683	BE264633	Hs.143638	WD repeat domain 4
	103723 103723	BE274312	Hs.214783	Homo saplens cDNA FLJ14041 fis, clone HE
	133260 133260	AA403045	Hs.6906	Homo sapiens cDNA: FLJ23197 fis, clone R
50	103766 103766	AI920783	Hs.191435	ESTs
	132051 132051	AA393968	Hs.180145	HSPC030 protein
	135289 135289	AW372569	Hs.9788	hypothetical protein MGC10924 similar to
	103794 103794	AF244135	Hs.30670	hepatocellular carcinoma-associated anti
	134319 134319	BE304999	Hs.285754	fumarate hydratase
55	119159 119159	AF142419	Hs.15020	homolog of mouse quaking QKI (KH domain
	103850 103850	AA187101	Hs.213194	hypothetical protein MGC10895
	322026 322026	AW024973	Hs.283675	NPD009 protein
	103861 103861	AA206236	Hs.4944	hypothetical protein FLJ12783
	447735 447735	AA775268	Hs.6127	Homo sapiens cDNA: FLJ23020 fis, clone L
60	131236 131236	AF043117	Hs.24594	ubiquitination factor E4B (homologous to
	129013 129013	AA371156	Hs.107942	DKFZP564M112 protein
	103988 103988	AA314389	Hs.342849	ADP-ribosylation factor-like 5
	425284 425284	AF155568	Hs.348043	NS1-associated protein 1
	133281 133281	AK001601	Hs.69594	high-mobility group 20A
65	108154 108154	NM_005754	Hs.220689	Ras-GTPase-activating protein SH3-domain
	135073 135073	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
	129593 129593	Al338247	Hs.98314	Homo saplens mRNA; cDNA DKFZp586L0120 (f
	132064 132064	AA121098	Hs.3838	serum-Inducible kinase
	131427 131427	AF151879	Hs.26706	CGI-121 protein
70	104282 104282	C14448	Hs.332338	EST
	130443 130443	D25216	Hs.155650	KIAA0014 gene product
	132837 132837	AA370362	Hs.57958	EGF-TM7-latrophilin-related protein
	104334 104334	D82614	Hs.78771	phosphoglycerate kinase 1
	134731 134731	D89377	Hs.89404	msh (Drosophila) homeo box homolog 2
75	131670 131670	H03514	Hs.15589	ESTs
	104402 104402	H56731	Hs.132956	ESTs

	120077 120077	17/79A	Un 400470	ECTo
	129077 129077 134927 134927	N74724 L36531	Hs.108479 Hs.91296	ESTs integrin, alpha 8
	134498 134498	AW246273	Hs.84131	threonyl-tRNA synthetase
	104488 104488	N56191	Hs.106511	protocadherin 17
5	129214 129214	AL044335	Hs.109526	zinc finger protein 198
	104530 104530	AK001676	Hs.12457	hypothetical protein FLJ10814
	104544 104544	Al091173	Hs.222362	ESTs, Weakly similar to p40 [H.sapiens]
	104567 104567	AA040620	Hs.5672	hypothetical protein AF140225
10	129575 129575	F08282	Hs.278428	progestin induced protein
10	104599 104599 104667 104667	AW815036 Al239923	Hs.151251 Hs.63931	ESTs ESTs
	104764 104764	AI039243	Hs.278585	ESTs
	104787 104787	AA027317	· IOLE I GOOG	gb:ze97d11.s1 Soares_fetal_heart_NbHH19W
	104804 104804	AJ858702	Hs.31803	ESTs, Weakly similar to N-WASP [H.saplen
15	130828 130828	AW631469	Hs.203213	ESTs
	104943 104943	AF072873	Hs.114218	frizzled (Drosophila) homolog 6
	105024 105024	AA126311	Hs.9879	ESTs
	105038 105038	AW503733	Hs.9414	KIAA1488 protein
20	105096 105096	AL042506	Hs.21599	Kruppel-like factor 7 (ubiquitous)
20	105169 105169 130401 130401	BE245294 BE396283	Hs.180789 Hs.173987	S164 protein eukaryotic translation initiation factor
	130114 130114	AA233393	Hs.14992	hypothetical protein FLJ11151
	105337 105337	Al468789	Hs.347187	myotubularin related protein 1
	105376 105376	AW994032	Hs.8768	hypothetical protein FLJ10849
25	131962 131962	AK000046	Hs.343877	hypothetical protein FLJ20039
	128658 128658	BE397354	Hs.324830	diptheria toxin resistance protein requi
	105508 105508	AA173942	Hs.326416	Homo sapiens mRNA; cDNA DKFZp564H1916 (f
	135172 135172	AB028956 AL137751	Hs.12144	KIAA1033 protein Homo sapiens mRNA; cDNA DKFZp434l0812 (f
30	132542 132542 105659 105659	AA283044	Hs.263671 Hs.25625	hypothetical protein FLJ11323
50	105674 105674	AI609530	Hs.279789	histone deacetylase 3
	105722 105722	Al922821	Hs.32433	ESTs
	115951 115951	BE546245	Hs.301048	sec13-like protein
25	105985 105985	AA406610		gb:zv15b10.s1 Soares_NhHMPu_S1 Homo sapi
35	131216 131216	AI815486	Hs.243901	Homo sapiens dDNA FLJ20738 fis, clone HE
	113689 113689	AB037850	Hs.16621	DKFZP434I116 protein
	130839 130839 130777 130777	AB011169 AW135049	Hs.20141 Hs.26285	similar to S. cerevisiae SSM4 Homo sapiens cDNA FLJ10643 fis, clone NT
	106196 106196	AA525993	Hs.173699	ESTs, Weakly similar to ALU1_HUMAN ALU S
40	133200 133200	AB037715	Hs.183639	hypothetical protein FLJ10210
	106328 106328	AL079559	Hs.28020	KIAA0766 gene product
	106423 106423	AB020722	Hs.16714	Rho guanine exchange factor (GEF) 15
	439608 439608	AW864696	Hs.301732	hypothetical protein MGC5306
45	106503 106503 106543 106543	AB033042 AA676939	Hs.29679 Hs.69285	cofactor required for Sp1 transcriptiona neuropilin 1
73	106589 106589	AK000933	Hs.28661	Homo sapiens cONA FLJ10071 fis, done HE
	106596 106596	AA452379		ESTs, Moderately similar to ALU7_HUMAN A
	106636 106636	AW958037	Hs.286	ribosomal protein L4
	131353 131353	AW754182		gb:RC2-CT0321-131199-011-c01 CT0321 Homo
50	131710 131710	NM_015368	Hs.30985	pannexin 1
	131775 131775	AB014548	Hs.31921	KIAA0648 protein
	106773 106773 106817 106817	AA478109 D61216	Hs.188833 Hs.18672	ESTs ESTs
	106848 106848	AA449014	Hs.121025	chromosome 11 open reading frame 5
55	418699 418699	BE539639	Hs.173030	ESTs, Weakly similar to ALU8_HUMAN ALU S
	130638 130638	AW021276	Hs.17121	ESTs
	107059 107059	BE614410	Hs.23044	RAD51 (S. cerevistae) homolog (E coli Re
	107115 107115	BE379623	Hs.27693	peptidylprolyl isomerase (cyclophilin)-l
60	107156 107156	AA137043	Hs.9663	programmed cell death 6-interacting prot
60	130621 130621 132626 132626	AW513087 AW504732	Hs.16803 Hs.21275	LUC7 (S. cerevisiae)-like hypothetical protein FLJ11011
	131610 131610	AA357879	Hs.29423	scavenger receptor with C-type lectin
	107295 107295	AA186629	Hs.80120	UDP-N-acetyl-elpha-D-galactosamine:polyp
	107315 107315	AA316241	Hs.90691	nudeophosmin/nudeoplasmin 3
65	107328 107328	AW959891	Hs.76591	KIAA0887 protein
	134715 134715	U48263	Hs.89040	prepronociceptin
	129938 129938	AW003668	Hs.135587	Human clone 23629 mRNA sequence
	130074 130074	AL038596	Hs.250745	polymerase (RNA) III (DNA directed) (62k
70	132036 132036 113857 113857	AL157433 AW243158	Hs.37706 Hs.5297	hypothetical protein DKFZp434E2220 DKFZP564A2416 protein
, 0	130419 130419	AF037448	Hs.155489	NS1-associated protein 1
	132616 132616	BE262677	Hs.283558	hypothetical protein PRO1855
	132358 132358	NM_003542	Hs.46423	H4 histone family, member G
	125827 125827	NM_003403	Hs.97496	YY1 transcription factor
75	107609 107609	R75654	Hs.164797	hypothetical protein FLJ13693
	107714 107714	AA015761	Hs.60642	ESTs

	407000 407000	4 4 0004 4770		about Coast at Coast with MMUD Home
	107832 107832	AA021473	11. 004004	gb:ze66c11.s1 Soares refina N2b4HR Homo
	124337 124337	N23541	Hs.281561	Homo saplens dDNA: FLJ23582 fis, clone L
	129577 129577	N75346	Hs.306121	CDC20 (cell division cycle 20, S. cerevi
_	132000 132000	AW247017	Hs.36978	melanoma antigen, family A, 3
5	107935 107935	AA029428	Hs.61555	ESTs
	131461 131461	AA992841	Hs.27263	KIAA1458 protein
	108029 108029	AA040740	Hs.62007	ESTs
	108084 108084	AA058944	Hs.116602	Homo saptens, clone IMAGE:4154008, mRNA,
	108168 108168	Al453137	Hs.63176	ESTs
10	108189 108189	AW376061	Hs.63335	ESTs, Moderately similar to A46010 X-lin
	108203 108203	AW847814	Hs.289005	Homo saptens cDNA: FLJ21532 fis, clone C
	108217 108217	AA058686	Hs.62588	ESTs
	108277 108277	AA064859		gb:zm50f03.s1 Stratagene fibroblast (937
	108309 108309	AA069818		gb:zm67e03.r1 Stratagene neuroepithelium
15	108340 108340	AA069820	Hs.180909	peroxiredoxin 1
	108427 108427	AA076382		gb:zm91g08.s1 Stratagene ovartan cancer
	108439 108439	AA078986		gb:zm92h01.s1 Stratagene ovarian cancer
	108469 108469	AA079487		gb:zm97f08.s1 Stratagene colon HT29 (937
	108501 108501	AA083256		gb:zn08g12.s1 Stratagene hNT neuron (937
20	108562 108562	AA100796		gb:zm26c06.s1 Stratagene pancreas (93720
	130890 130890	Al907537	Hs.76698	stress-associated endoplasmic reticulum
	130385 130385	AW067800	Hs.155223	stanniocalcin 2
	108807 108807	Al652236	Hs.49376	hypothetical protein FLJ20644
	108833 108833	AF188527	Hs.61661	ESTs, Weakly similar to AF174605 1 F-box
25	108846 108846	AL117452	Hs.44155	DKFZP586G1517 protein
	131474 131474	L46353	Hs.2726	high-mobility group (nonhistone chromoso
	108941 108941	AA148650		gb:zo09e06.s1 Stratagene neuroepithelium
	108996 108996	AW995610	Hs.332436	EST
	131183 131183	Al611807	Hs.285107	hypothetical protein FLJ13397
30	109022 109022	AA157291	Hs.21479	ubinuclein 1
	109068 109068	AA164293	Hs.72545	ESTs
	129021 129021	AL044675	Hs.173081	KIAA0530 protein
	109146 109146	AA176589	Hs.142078	EST
	131080 131080		Hs.2271	endothelin 1
35	109222 109222	AA192833	Hs.333512	similar to rat myomegalin
	109481 109481	AA878923	Hs.289069	hypothetical protein FLJ21016
	109516 109516		Hs.71913	ESTs
	109556 109556		Hs.87385	ESTs
	109578 109578		Hs.27214	ESTs
40	109625 109625		Hs.22697	ESTs
	109648 109648		Hs.7154	ESTs
	109699 109699		Hs.167483	ESTs
	109933 109933		Hs.20945	Homo sapiens clone 24993 mRNA sequence
	110039 110039	H11938	Hs.21907	histone acetyttransferase

TABLE 2A

Table 2A shows the accession numbers for those pkeys lacking unigenelD's for Table 2. The pkeys in Table 7 lacking unigenelD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

 $\begin{array}{ccc} 10 & \text{Pkey:} & \text{Unique Eos probeset identifier number} \\ & \text{CAT number:} & \text{Gene cluster number} \end{array}$

Accession: Genbank accession numbers

15	Pkey	CAT Number	Accession
	108469 108501	116761_1 1368412	AA079487 AA128547 AA128291 AA079587 AA079600 AA083256
20	108562 101300	36375_1 4669_1	AA100796 AF020589 AA074629 AA075946 AA100849 AA085347 AA126309 AA079311 AA079323 AA085274 BE535511 M62098 AA306787 AW891766 AA348998 AA338869 AA344013 AW956561 AW389343 AW403607 L40391 AW408435 AA121738 AI568978 H13317 R20373 AW948724 AW948744 AA335023 AA436722 AA448690 C21404 AW884390 AA345454 AA303292 AA174174 BE092290 T90614 AA035104 R76028 AA126924 AA741086 AW022056 AW118940 AA121666 AI832409 AA683475 AI140901 AI623576 AW519064 AW474125 AI953923 AI735349 AW150109 AI438154 AW118130 AW270782 AI804073 N27434 AA876543 AA937815 AI051166 AA505378 AI041975 AI335355
25			AI089540 AA662243 AI127912 AI925604 AI250880 AI366874 AI554386 AI815196 AI683526 AI435885 AI160934 H79030 AI801493 AA448691 AI673767 AI076042 AI804327 AA813438 AA680002 AI274492 T16177 AI287337 AI935050 AA907805 AA911493 AI589411 AI371358 AW576236 AI078866 AW516168 AA346372 AI560185 AA471009 R75857 AA296025 AA523155 AA853168 AI696593 AI658482 AI566601 AW072797 AA128047 AA035502 AW243274 AA992517 R43760
30	117156 125565	145392_1 1704098_1	W73853 AA928112 W77887 AW889237 AA148524 AI749182 AI754442 AI338392 AI253102 AI079403 AI370541 AI697341 H97538 AW188021 AI927669 W72716 AI051402 AI188071 AI335900 N21488 AW770478 W92522 AI691028 AI913512 AI144448 W73819 AA604358 N28900 W95221 AI868132 H98465 AA148793 R20840 R20839
35	132983	11922_1	M30269 NM_002508 X82245 Al078760 AW957003 D78945 M27445 AA650439 AL048816 AV660256 AV660347 AA333052 BE295257 T60999 AA383049 AW369677 Z26985 AW175704 AA343326 AW747957 Al818389 W17308 W17302 H15591 AA371284 AA370412 W94966 BE384365 T28498 R80714 R16959 H21723 AW835154 D56097 D56381 W21232 AA190565 AW379755 AW057895
40	133681	13893_1	Al352558 Z82248 X78138 NM_003405 AU077248 AA223125 S80794 D78577 Al124697 AW403970 BE614089 BE296713 BE621334 L20422 X80536 D54224 D54950 X57345 N29226 AA127798 AA340253 F08031 AA192540 H67636 AA321827 AW950283 AA084159 BE538808 AW401377 AA256774 C03366 W46595 W47608 AA305009 H69431 H69456 AL120082 H11706 AA303717 AA361357 H22042 H78020 AW999584 AA134368 AA322911 AA322961 H60980 N85248 N31547 H79624 T11718 W85826 AW894663 AW894624 BE167441 BE170015 AA304626 AW602163 AW998929 AA156681
45			AA151067 BE002724 AA608688 H82692 BE155392 AW383636 BE155394 AA487004 AW383504 AI342365 R82553 W16498 BE155344 AI143938 R69901 AA322873 AW340648 R25364 AA367935 AI559406 AA033522 AA374252 AW835019 AI922133 AI697089 N99662 AW189078 AI199076 AW151589 W59944 AA662875 W94022 AA299055 AI039008 AI829449 AA583503 AI635674 AW131665 AI473820 AW273118 AW900930 AA908944 AI688035 AW170272 AI082545 AW468176 AI608761 AI082748 AI911682 AI248943 AI831016 AA192465 AI218477 AA938406 AA385288 AI809817 AA905196 AI191245 AI470204 AI188296 AI421367 AI125315 AI087141 AA629032 AA740589 AI554181
50			AA150830 Al248541 Al077943 AA775958 AA864930 Al261476 Al123121 Al310394 AA862331 AA872478 BE537084 Al205606 AA720684 Al872093 AW150042 AL120538 AA219627 AA988608 C21397 Al359337 H25337 Al089749 AA605146 Al359620 AA150478 Al359738 AW383642 AW995424 Al766457 R56892 Al089839 W61343 N69107 W46459 AA565955 N20527 Al279782 W46596 AA776573 H23204 Al866231 Al083995 N21530 AA126874 D82630 W65437 Al086917 AW382095 Al086877 H69844 AW340217 W85827 L08439 AA262704 AA565380 W47413 W94135 AA223241
55			AW089153 AA084101 BE538000 AA096126 T28031 AA491574 R84813 AA774536 AW383522 AA155615 AW383529 AA491520 AW028427 AA171496 AJ469689 AW664539 AI811102 AI811116 BE464590 BE350791 H78021 T15405 H21979 AA219489 H13301 AA505883 AI864305 AJ423963 AW084401 F04963 R69858 H67097 AJ917740 AJ655561 H69864 AA033631 AW383484 AJ886261 H25293 AA513281 AW271187 H11617 N79982 AJ174338 AJ904207 AJ904208 BE614558 W94127 W65436 AJ272249 AA700018 AJ579932 AJ085941 AW152629
60	121335 130018 121822	279548_1 18986_1 244391_1	AA404418 AI217248 AA53093 AW957317 AW872498 AI560785 AI289110 AW135512 X97261 T68873 AI743860 N49543 AW027759 BE349467 AI656284 BE463975 R35022 AA370031 AW955302 AL042109 N53092 AI611424 AL079362 AI969290 AI928016 BE394912 BE504220 BE467505 AI611611 AI611407 AI611452 W56437 AI284566 AL079362 AI969290 AI928016 BE394912 BE504220 BE467505 AI611611 AI611407 AI611452 W56437 AI284566 AL079362 AI969290 AI928058 AI308085 AI074952 AA437315 AA628161 AW301728 AI150224 AA400137 AA437279 AI223355 AA639462 AI261373 AI432414 AI984994 AI539335 AA401550 AA358757 AI609976 AA442357 AA359393 AA437046
65			AA370301 AA429328 AW272055 AI580502 AI832944 AI038530 AA425107 AI014986 AI148349 AW237721 AW779756 AW137877 AI125293 AA400404 R28554
70	108309 107832 123523 123964 118475 104787	111495_1 genbank_AA021 genbank_AA608 genbank_C1398 genbank_N6684 genbank_AA027	3588 AA608588 61 C13961 15 N66845
75	106596 113947 108277	304084_1 genbank_W8474 genbank_AA064	Al583948 AA578212 AW303715 AA653450 AA456981 Al400385 W88533 Al224133 AW272145 AA088686 R94698 68 W84768

	108427	genbank_AA076382	AA076382
	108439	genbank_AA078986	AA078986
	131353	231290_1 AW411259 H235	55 AW015049 Al684275 AW015886 AW068953 AW014085 Al027260 R52686 AA918278 Al129462
_		AA969360 N3486	59 A1948416 AA534205 AA702483 AA705292
5	101447	entrez_M21305 M21305	
	108931	genbank_AA147186	AA147186
	108941	genbank_AA148650	AA148650
	103138	entrez_X65965 X65965	•
	119174	genbank_R71234 R71234	
10	119416	genbank_T97186 T97186	
	105985	genbank_AA406610	AA406610
	100327	entrez_D55640 D55640	•

TABLE 3:

5

Pkey: Accession: ExAccn: UnigeneID: Unigene Title: Unique Eos probeset identifier number Accession number used for previous patent filings Exemplar Accession number, Genbank accession number Unigene number Unigene gene title

10					
	Pkey	Accession	ExAccn	UniGene	UnigeneTitle
		D86425	AW291587		nidogen 2
1.5		D86983	D86983	Hs.118893	Melanoma associated gene
15		HG1098-HT1098 HG1103-HT1103		Hs.121489	cystatin D v-ral simian leukemia viral oncogene hom
		HG3342-HT3519	_		Inhibitor of DNA binding 1, dominant neg
	100991		J03836	Hs.82085	serine (or cysteine) proteinase inhibito
		L06797	BE245301		chemokine (C-X-C motif), receptor 4 (fus
20		L15388		3Hs.211569	G protein-coupled receptor kinase 5
	101194	L20971	L20971	Hs.188	phosphodiesterase 4B, cAMP-specific (dun
		L35545	D30857	Hs.82353	protein C receptor, endothelial (EPCR)
		L76380		5Hs.152175	calcitonin receptor-like
25		M21305 M24736	M21305 AA296520	LIA GOEAG	gb:Human alpha satellite and satellite 3 selectin E (endothelial adhesion molecul
23		M31166	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
		M31551	Y00630	Hs.75716	serine (or cysteine) proteinase inhibito
		M32334		Hs.347326	Intercellular adhesion molecule 2
	101674	M61916	NM_00229	1Hs.82124	laminin, beta 1
30	101714	M68874	M68874	Hs.211587	phospholipase A2, group IVA (cytosolic,
		M74719		9Hs.326198	transcription factor 4
		M92934	BE243845		connective tissue growth factor
		M94856 U03057		Hs.153179 Hs.118400	fatty acid binding protein 5 (psortasis- singed (Drosophila)-like (sea urchin fas
35		U03877	AA301867		EGF-containing fibulin-like extracellula
55		U18300	NM_00010		damage-specific DNA binding protein 2 (4
	102241	U27109		1Hs.268107	multimerin
	102283	U31384	AW161552	Hs.83381	guantne nucleotide binding protein 11
40		U33053	U33053	Hs.2499	protein kinase C-like 1
40		U59423	U59423	Hs.79067	MAD (mothers against decapentaplegic, Dr
		U70322 U81607	NM_00510	0Hs.168075	karyopherin (importin) beta 2 A kinase (PRKA) anchor protein (gravin)
		U83463	AF000652		syndecan binding protein (syntenin)
		U89942	NM_00231		lysyl oxidase-like 2
45		X04729	J03836	Hs.82085	serine (or cysteine) proteinase inhibito
	102898	X06256	NM_00220	5Hs.149609	Integrin, alpha 5 (fibronectin receptor,
		X07820	X07820	Hs.2258	matrix metalloproteinase 10 (stromelysin
		X54925	M13509	Hs.83169	matrix metalloproteinase 1 (interstifial
50		X54936 X60957	BE018302 NM_00542		placental growth factor, vascular endoth tyrosine kinase with immunoglobulin and
50		X67235	-	Hs.118651	hematopoietically expressed homeobox
		X67951		Hs.180909	peroxiredoxin 1
•		X69910	NM_00682		transmembrane protein (63kD), endoplasmi
	103280	X79981	U84722	Hs.76206	cadherin 5, type 2, VE-cadherin (vascula
55		Z18951	A1878826	Hs.74034	caveolin 1, caveolae protein, 22kD
		AA187101		Hs.213194	hypothetical protein MGC10895
		N24990	Z44203 AW630488	Hs.26418	ESTs
		R81003 AA025351	AVV030400 AI039243		protease, serine, 23 ESTs
60		AA027168	AA027167		KIAA0955 protein
		AA040465	AL133035		hypothetical protein DKFZp434G171
		AA045136	T79340	Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
		AA054087	AF065214		phospholipase A2, group IVC (cytosolic,
CE		AA071089		Hs.345588	desmoplakin (DPI, DPII)
65		AA085918	Y12059 AA313825	Hs.278675	bromodomain-containing 4 AD036 protein
		AA187490 AA227926	AW388633		solute carrier family 7, (cationic amino
	105233	AA234743	AW338625		ESTs
	105376	AA236559	AW994032		hypothetical protein FLJ 10849
70	105729	AA292694	H46612	Hs.293815	Homo sapiens HSPC285 mRNA, partial cds
	105826	AA398243		Hs.194477	E3 ubiquitin ligase SMURF2
	105977	AA406363		Hs.30822	hypothetical protein FLJ11110
		AA411465	AB033888		SRY (sex determining region Y)-box 18
75		AA412284	X64116 H93366	Hs.171844 Hs.7567	Homo sapiens cDNA: FLJ22296 fis, clone H Homo sapiens cDNA: FLJ21962 fis, clone H
13	100124	AA423987	กรวว00	(10.730)	HUMBU SAPIEMS WHAT PLUZ 130Z HS, CIONE M

	100166 AAA06000	AA425414 Hs.33287	avalant factor I/D
	106155 AA425309 106302 AA435896	AA398859 Hs.18397	nuclear factor VB
	106423 AA448238	AB020722 Hs.16714	hypothetical protein FLJ23221 Rho guanine exchange factor (GEF) 15
	106793 AA478778	H94997 Hs.16450	ESTs
5	107174 AA621714	BE122762 Hs.25338	ESTS
3	107216 D51069	D51069 Hs.211579	metanoma cell adhesion molecule
	107295 T34527	AA186629 Hs.80120	UDP-N-acetyl-alpha-D-galactosamine:polyp
	107385 U97519	NM_005397Hs.16426	podocalyxin-like
	108756 AA127221	AA127221 Hs.117037	ESTs
10	108846 AA132983	AL117452 Hs.44155	DKFZP586G1517 protein
10	108888 AA135606	AA135606 Hs.189384	gb:zi10a05.s1 Soares_pregnant_uterus_NbH
	109001 AA156125	Al056548 Hs.72116	hypothetical protein FLJ20992 similar to
	109166 AA179845	AA219691 Hs.73625	RAB6 interacting, kinesin-like (rabkines
	109456 AA232645	AW956580 Hs.42699	ESTs
15	109768 F10399	F06838 Hs.14763	ESTs
13	110107 H16772	AW151660 Hs.31444	ESTS
	110906 N39584	AA035211 Hs.17404	ESTs
	110984 N52006	AW613287 Hs.80120	UDP-N-acetyl-alpha-D-galactosamine:polyp
	111006 N53375	BE387014 Hs.166146	Homer, neuronal immediate early gene, 3
20	111018 N54067	Al287912 Hs.3628	mitogen-activated protein kinase kinase
20	111133 N64436	AW580939 Hs.97199	complement component C1q receptor
	111760 R26892	BE551929 Hs.268754	Homo sapiens cDNA FLJ11949 fis, clone HE
	113073 T33637	N39342 Hs.103042	microtubule-associated protein 1B
	113195 T57112	H83265 Hs.8881	ESTs, Weakly similar to S41044 chromosom
25	113923 W80763	AW953484 Hs.3849	hypothetical protein FLJ22041 similar to
23	114521 AA046808	AW139036 Hs.108957	40S ribosomal protein S27 isoform
	115061 AA253217	AI751438 Hs.41271	Homo sapiens mRNA full length insert cDN
	115096 AA255991	Al683069 Hs.175319	ESTs
	115145 AA258138	AA740907 Hs.88297	ESTs
30	115819 AA426573	AA486620 Hs.41135	endomucin-2
50	115947 AA443793	R47479 Hs.94761	KIAA1691 protein
	116314 AA490588	AI799104 Hs.178705	Homo sapiens cONA FLJ11333 fis, clone PL
	116339 AA496257	AK000290 Hs.44033	dipeptidyl peptidase 8
	116430 AA609717	AK001531 Hs.66048	hypothetical protein FLJ10669
35	116589 D59570	Al557212 Hs.17132	ESTs, Moderately similar to I54374 gene
-	116733 F13787	AL157424 Hs.61289	synaptojanin 2
	117023 H88157	AW070211 Hs.102415	Homo sapiens mRNA; cDNA DKFZp586N0121 (f
	117186 H98988	H98988 Hs.42612	ESTs, Weakly similar to ALU1_HUMAN ALU S
	117563 N34287	AF055634 Hs.44553	unc5 (C.elegans homolog) c
40	117997 N52090	N52090 Hs.47420	EST
	118475 N66845	N66845	gb:za46c11.s1 Soares fetal liver spleen
	118581 N68905	N68905	gb:za69b09.s1 Soares_fetal_lung_NbHL19W
	119073 R32894	BE245360 Hs.279477	ESTs
	119155 R61715	R61715 Hs.310598	ESTs, Moderately similar to ALU1_HUMAN A
45	119174 R71234	R71234	gb:yi54c08.s1 Soares placenta Nb2HP Homo
	119221 R98105	C14322 Hs.250700	tryptase beta 1
	119416 T97186	T97186	gb:ye50h09.s1 Soares fetal liver spleen
	119866 W80814	AA496205 Hs.193700	Homo sapiens mRNA; cDNA DKFZp58610324 (f
	121335 AA404418	AA404418	gb:zw37e02.s1 Soares_total_fetus_Nb2HF8_
50	121381 AA405747	AW088642 Hs.97984	hypothetical protein FLJ22252 similar to
	123160 AA488687	AA488687 Hs.284235	ESTs, Wealty similar to 138022 hypotheti
	123473 AA599143	AA599143	gb:ae52d04.s1 Stratagene tung carcinoma
	123523 AA608588	AA608588	gb:ae54e06.s1 Stratagene lung carcinoma
	123533 AA608751	AA608751	gb:ae56h07.s1 Stratagene lung carcinoma
55	123964 C13961	C13961	gb:C13961 Clontech human aorta polyA+mR
	124006 D60302	Al147155 Hs.270016	ESTs
	124315 H94892	NM_005402Hs.288757	v-ral simian leukemia viral oncogene hom
	124659 N93521	Al680737 Hs.289068	Homo sapiens cDNA FLJ11918 fis, clone HE
	124669 N95477	Al571594 Hs.102943	hypothetical protein MGC12916
60	124847 R60044	W07701 Hs.304177	Homo sapiens done FLB8503 PRO2286 mRNA,
	124875 R70506	Al887664 Hs.285814	sprouty (Drosophila) homolog 4
	125091 T91518	T91518	gb:ye20f05.s1 Stratagene lung (937210) H
	125103 T95333	AA570056 Hs.122730	ESTs, Moderately similar to KIAA1215 pro
	125355 R45630	R60547 Hs.170098	KIAA0372 gene product
65	125565 R20839	R20840	gb:yg05c08.r1 Soares infant brain 1NIB H
	125590 . R23858	R23858 Hs.143375	Homo sapiens, clone IMAGE:3840937, mRNA,
	126511 Al024874	T92143 Hs.57958	EGF-TM7-latrophilin-related protein
	126563 W26247	AA516391 Hs.181368	U5 snRNP-specific protein (220 kD), orth
	126649 AA856990	AA001860 Hs.279531	ESTs
70	126872 AA136653	AW450979	gb:UI-H-BI3-ala-a-12-0-UI.s1 NCI_CGAP_Su
	127402 AA358869	AA358869 Hs.227949	SEC13 (S. cerevisiae)-like 1
	127651 AJ123976	AA382523 Hs.105689	MSTP031 protein
	127759 Al369384	Al369384 Hs.292441	ESTs
~~	128062 AA379500	AA379621 Hs.105547	neural proliferation, differentiation an
75	128992 R49693	H04150 Hs.107708	ESTs
	129046 AA195678	AB029290 Hs.108258	actin binding protein; macrophin (microf

	129188	M30257	NM 00107	'8Hs.109225	vascular cell adhesion molecule 1
	129314	AA028131		Hs.290356	mesodem development candidate 1
	129371	M10321	X06828	Hs.110802	von Willebrand factor
_	129468	J03040		Hs.111779	secreted protein, acidic, cysteine-rich
5	129765	M86933	M86933	Hs.1238	amelogenin (Y chromosome)
		AA012933	AA012848	Hs.12570	tubulin-specific chaperone d
		AA286710	AF055581		lysosomal
		AA243278		Hs.109059	mitochondrial ribosomal protein L12
10		D59711		Hs.17132	ESTs, Moderately similar to 154374 gene
10		T94452 AA053400		Hs.201591	ESTs
		AA370302	D81866	Hs.203213 Hs.21739	ESTs Homo sapiens mRNA; cDNA DKFZp586I1518 (f
		J05008	NM_00195		endothelin 1
		U85193	W27392	Hs.33287	nuclear factor I/B
15		AA256153	AI824144		ESTs
	131486	X83107	F06972	Hs.27372	BMX non-receptor tyrosine kinase
		AA046593	AA040311	Hs.28959	ESTs
		AA410480	AA359615		ESTs
20		D45304	AA443966		ESTs
20		M90657	AW960564		transmembrane 4 superfamily member 1
		AA010163 AA136353	AW361018 Al267615		upstream regulatory element binding prot ESTs
		Y07867		Hs.38022 Hs.279663	Pirin
		U84573	AI752235		procollagen-lysine, 2-oxoglutarate 5-dio
25		X60486	NM_00354		H4 histone family, member G
		AA132969		Hs.260116	metalloprotease 1 (pitrilysin family)
	132456	AA114250	AB011084	Hs.48924	KIAA0512 gene product; ALEX2
		F13782	NM_00129	0Hs.4980	LIM domain binding 2
20		AA283035	N92589	Hs.261038	ESTs, Wealdy similar to 138022 hypotheti
30		AB002301	AB002301		KIAA0303 protein
		AA056731	NM_00460		Sjogren syndrome antigen A2 (60kD, ribon
		U68019 H99198	AWU81883 AA125985	Hs.211578	Homo saplens cDNA: FLJ23037 fis, clone L
		AA598702	BE263252		thymosin, beta, identified in neuroblast hypothetical protein MGC3178
35		N77151	AF234532		myosin X
-		AA505133		Hs.279905	clone HQ0310 PRO0310p1
	133061	AB000584	Al186431	Hs.296638	prostate differentiation factor
		D12763	AA026533		interleukin 1 receptor-like 1
40		AA253193	AW021103		hypothetical protein FLJ20373
40		AA432248		Hs.183639	hypothetical protein FLJ10210
		AA083572 AA479713	AA403045 A1866286	Hs.71962	Homo sapiens cDNA: FLJ23197 fis, clone R
		L40395	BE619053		ESTs, Weakly similar to B36298 proline-r eukaryotic translation initiation factor
		X52947	NM_00016		gap Junction protein, alpha 1, 43kD (con
45		W80846	Al129903	Hs.74669	vesicle-associated membrane protein 5 (m
	133607	M34539	BE273749	•	FK506-binding protein 1A (12kD)
		D67029	NM_003003		SEC14 (S. cerevisiae)-like 1
		U09587	NM_002047		glycyl-tRNA synthetase
50		M85289	M85289	Hs.211573	heparan sulfate proteoglycan 2 (pertecan
50		D10522 W84712	AI878921 AU076964	Hs.75607	myristoylated alanine-rich protein kinas
		D29992	C18356	Hs.295944	calumenin tissue factor pathway inhibitor 2
		L34657	L34657	Hs.78146	platelet/endothelial cell adhesion molec
		S78569	NM_002290		laminin, alpha 4
55	134088	D43636	Al379954	Hs.79025	KIAA0096 protein
	134161	U97188	AA634543		IGF-II mRNA-binding protein 3
		AA487558	AW580939		complement component C1q receptor
		M28882	X68264	Hs.211579	melanoma cell adhesion molecule
60		X70683		Hs.83484	SRY (sex determining region Y)-box 4
00		X14787 AA236324	AI750878 AW968058		thrombospondin 1 nudix (nucleoside diphosphate linked moi
		C15324	Al272141	Hs.83484	SRY (sex determining region Y)-box 4
		AA452000	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
		D83174	AA114212	Hs.9930	serine (or cysteine) proteinase inhibito
65		D00596	X02308	Hs.82962	thymidylate synthetase
		D11428	NM_000304		peripheral myelin protein 22
		D13640	AU076465		KIAA0015 gene product
		D14874	H73444	Hs.394	adrenomedullin
70		D26129 D28476	NM_002933 AL121516		ribonuclease, RNase A family, 1 (pancrea
, ,		D86425	AW291587		thyroid hormone receptor interactor 12 nidogen 2
		D86983	D86983	Hs.118893	Melanoma associated gene
		D87953	AW888941		N-myc downstream regulated
	100529	HG1862-HT1897	BE313693		calmodulin 2 (phosphorylase kinase, delt
75		HG2614-HT2710		Hs.114599	collagen, type VIII, atpha 1
	100619	HG2639-HT2735	N24433	Hs.241567	RNA binding motif, single stranded inter

		HG2855-HT2995		Hs.180414	heat shock 70kD protein 2
		HG3044-HT3742		Hs.287820	fibronectin 1
		HG3342-HT3519 HG3543-HT3739		Hs.75424	inhibitor of DNA binding 1, dominant neg
5		HG4069-HT4339		Hs.303649	insulin-like growth factor 2 (somatomed) small inducible cytokine A2 (monocyte ch
		HG417-HT417		Hs.297939	cathepsin B
	100991		J03836	Hs.82085	serine (or cysteine) proteinase inhibito
	101097	L06797 L08246	BE245301 Al439011	Hs.89414 Hs.86386	chemokine (C-X-C motif), receptor 4 (fus
10	101142		L12711	Hs.89643	myeloid cell leukemia sequence 1 (BCL2-r transketolase (Wernicke-Korsakoff syndro
	101156		AA340987		prolylcarboxypeptidase (angiotensinase C
		L15388		8Hs.211569	G protein-coupled receptor kinase 5
	101184 101192		NM_001674 BE247295		activating transcription factor 3 solute carrier family 20 (phosphate tran
15	101317		L42176	Hs.8302	four and a half LIM domains 2
	101336		NM_006732		FBJ murine osteosarcoma viral oncogene h
	101345		_	5Hs.152175	calcitonin receptor-like
		M15990 M23254	M15990 BE410405	Hs.194148 Hs.76288	v-yes-1 Yamaguchi sarcoma viral oncogene calpain 2, (m/li) large subunit
20		M24736	AA296520		selectin E (endothelial adhesion molecul
		M26576	X12784	Hs.119129	collagen, type IV, alpha 1
		M27396	AA307680		asparagine synthetase
		M31166 M31994	M31166 BE293116	Hs.2050 Hs 76392	pentaxin-related gene, rapidly induced b aldehyde dehydrogenase 1 family, member
25		M32334		Hs.347326	intercellular adhesion molecule 2
		M35878	Al752416	Hs.77326	insulin-like growth factor binding prote
		M36429 M57730	AF064853		guanine nucleotide binding protein (G pr
		M57731	NM_004428 AV650262		ephrin-A1 GRO2 oncogene
30	101667		NM_005381		nucleolin
	101682		AF043045		filamin B, beta (actin-binding protein-2
	101714 101720		M68874 M69043	Hs.211587 Hs.81328	phospholipase A2, group IVA (cytosolic, nuclear factor of kappa light polypeptid
	101741		NM_003199		transcription factor 4
35	101744			Hs.118625	hexokinase 1
	101793 101837		W01076 M92843	Hs.278573 Hs.343586	CD59 antigen p18-20 (antigen identified
	101838		BE243845		zinc finger protein homologous to Zfp-36 connective tissue growth factor
40	101840		AA236291	Hs.183583	serine (or cysteine) proteinase inhibito
40	101857 101864		BE550723 BE392588	Hs.153179	fatty acid binding protein 5 (psoriasis-
	101931		NM_006823		transgelin protein kinase (cAMP-dependent, catalyti
	101966	S81914	X96438	Hs.76095	Immediate early response 3
45	102012 102013		BE259035		singed (Drosophila)-like (sea urchin fas
73	102013		BE616287 AA301867		catenin (cadherin-associated protein), a EGF-containing fibulin-like extracellula
	102059			Hs.76669	nicotinamide N-methyltransferase
	102121		NM_004998		myosin iE
50	102283 102300		AW161552 Al929721	Hs.83381 Hs.5120	guanine nucleotide binding protein 11 dynein, cytopiasmic, light polypeptide
50	102378		AU076887		spermidine/spermine N1-acetyltransferase
	102395	U41767	AU077005		a disintegrin and metalloproteinase doma
	102460			Hs.211582	myosin, light polypeptide kinase
55	102491 102499		U51010 BE243877	Hs.76941	gb:Human nicotinamide N-methyltransferas ATPase, Na+/K+ transporting, beta 3 poly
••	102523			Hs.15432	downregulated in ovarian cancer 1
	102560			Hs.63984	cadherin 13, H-cadherin (heart)
	102564 102589		U59423 AU076728	Hs.79067	MAD (mothers against decapentaplegic, Dr cysteine-rich, angiogenic inducer, 61
60	102600			Hs.66713	hepatitis delta antigen-interacting prot
	102645	U67963	AL119566		lysosomal
	102687		NM_007019		ubiquitin carrier protein E2-C
	102693 102709		AA532780 AA122237		eukaryotic translation initiation factor microsomal glutathione S-transferase 2
65	102759		NM_005100		A kinase (PRKA) anchor protein (gravin)
	102804		NM_002318		lysyl oxidase-like 2
	102882 102907		A1767736 BE409861	Hs.290070	gelsolin (amyloldosis, Finnish type)
	102915			Hs.2258	heme oxygenase (decycling) 1 matrix metalloproteinase 10 (stromelysin
70	102927	X12876	BE512730	Hs.65114	keratin 18
	102960			Hs.76053	DEAD/H (Asp-Gtu-Ala-Asp/His) box polypep
	103011 103020			Hs.326035 Hs.195464	early growth response 1 filamin A, alpha (actin-binding protein-
	103029		AW800726		GRO1 oncogene (melanoma growth stimulati
75	103036	X54925	M13509	Hs.83169	matrix metalloproteinase 1 (interstitial
	103056	X3/2U5	Y18024	Hs.78877	inosital 1,4,5-trisphosphate 3-kinase B

	402000 VE0700	A11077004 Na 00000	mulia D4 /DD4D4, constructed adapaments
	103080 X59798 103095 X60957	AU077231 Hs.82932 NM 005424Hs.78824	cyclin D1 (PRAD1: parathyroid adenomatos
	103138 X65965	X65965	tyrosine kinase with immunoglobulin and gb:H.sapiens SOD-2 gene for manganese su
	103176 X69111	AL021154 Hs.76884	inhibitor of DNA binding 3, dominant neg
5	103195 X70940	AA351647 Hs.2642	eukaryotic translation elongation factor
•	103347 X87838	AU077309 Hs.171271	catenin (cadherin-associated protein), b
	103371 X91247	X91247 Hs.13046	thioredoxin reductase 1
	103432 X97748	X97748	gb:H.sapiens PTX3 gene promotor region.
	103471 Y00815	Y00815 Hs.75216	protein tyrosine phosphatase, receptor t
10	103967 AA303711	AL120051 Hs.144700	ephrin-B1
	104447 L44538	AW204145 Hs.156044	ESTs
	104764 AA025351	Al039243 Hs.278585	ESTs
	104783 AA027050	AA533513 Hs.93659	protein disulfide isomerase related prot
1.5	104798 AA029462	AW952619 Hs.17235	Homo sapiens done TCCCIA00176 mRNA sequ
15	104865 AA045136	T79340 Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
	104877 AA047437 104894 AA054087	Al138635 Hs.22968 AF065214 Hs.18858	Homo saptens clone IMAGE:451939, mRNA se phospholipase A2, group IVC (cytosolic,
	104952 AA071089	AW076098 Hs.345588	desmoplakin (DPI, DPII)
	105113 AA156450	AB037816 Hs.8982	Homo sapiens, clone IMAGE:3506202, mRNA,
20	105178 AA187490	AA313825 Hs.21941	AD036 protein
	105196 AA195031	W84893 Hs.9305	angiotensin receptor-like 1
	105215 AA205724	AA205759 Hs.10119	hypothetical protein FLJ14957
	105263 AA227926	AW388633 Hs.6682	solute carrier family 7, (cationic amino
0.5	105271 AA227986	AA807881 Hs.25329	ESTs .
25	105330 AA234743	AW338625 Hs.22120	ESTs
	105461 AA253216	BE539071 Hs.69388	hypothetical protein FLJ20505
	105492 AA256210	Al805717 Hs.289112	CGI-43 protein
	105493 AA256268	AL047586 Hs.10283	RNA binding motif protein 8B
30	105594 AA279397 105727 AA292379	AB024334 Hs.25001 AL135159 Hs.20340	tyrosine 3-monooxygenase/tryptophan 5-mo KIAA1002 protein
50	105732 AA292717	AW504170 Hs.274344	hypothetical protein MGC12942
	105767 AA346551	AW370946 Hs.23457	ESTs
	105882 AA400292	W46802 Hs.81988	disabled (Drosophila) homolog 2 (mitogen
	105936 AA404338	Al678765 Hs.21812	ESTs
35	106031 AA412284	X64116 Hs.171844	Homo sapiens cDNA: FLJ22296 fis, clone H
	106124 AA423987	H93366 Hs.7567	Homo saplens cDNA: FLJ21962 fis, clone H
	106222 AA428594	AA356392 Hs.21321	Homo sapiens clone FLB9213 PRO2474 mRNA,
	106241 AA430108	BE019681 Hs.6019	Homo saplens cDNA: FLJ21288 fis, clone C
40	106263 AA431462	W21493 Hs.28329	hypothetical protein FLJ14005
40	106264 AA431470 106366 AA443756	AL046859 Hs.3407 AA186715 Hs.336429	protein kinase (cAMP-dependent, catalyti RIKEN cDNA 9130422N19 gene
	106454 AA449479	NM_014038Hs.5216	HSPC028 protein
	106634 AA459916	W25491 Hs.288909	hypothetical protein FLJ22471
	106724 AA465226	N48670 Hs.28631	Homo sapiens cDNA: FLJ22141 fis, clone H
45	106793 AA478778	H94997 Hs.16450	ESTs
	106799 AA479037	BE313412 Hs.7961	Homo sapiens done 25012 mRNA sequence
	106842 AA482597	AF124251 Hs.26054	novel SH2-containing protein 3
	106868 AA487561	BE185536 Hs.301183	molecule possessing ankyrin repeats indu
50	106890 AA489245 106961 AA504110	AA489245 Hs.88500	mitogen-activated protein kinase 8 inter
50	106974 AA520989	AW243614 Hs.18063 Al817130 Hs.9195	Homo sapiens cDNA FLJ10768 fis, clone NT Homo sapiens cDNA FLJ13698 fis, clone PL
	107030 AA599434	AL117424 Hs.25035	chloride intracellular channel 4
	107061 AA608649	BE147611 Hs.6354	stromal cell derived factor receptor 1
	107086 AA609519	NM_012331Hs.26458	methionine sulfoxide reductase A
55	107216 D51069	D51069 Hs.211579	melanoma cell adhesion molecule
	107385 U97519	NM_005397Hs.16426	podocalyxin-like
	107444 W28391	W28391 Hs.343258	proliferation-associated 2G4, 38kD
	107985 AA035638	T40064 Hs.71968	Homo sapiens mRNA; cDNA DKFZp564F053 (fr
60	108507 AA083514	AI554545 Hs.68301	ESTS
00	108695 AA121315 108931 AA147186	AB029000 Hs.70823 AA147186	KIAA1077 protein qb:zo38d01.s1 Stratagene endothelial cel
	109001 AA156125	AI056548 Hs.72116	hypothetical protein FLJ20992 similar to
	109195 AA188932	AF047033 Hs.132904	solute carrier family 4, sodium bicarbon
	109390 AA219653	AW007485 Hs.87125	EH-domain containing 3
65	109456 AA232645	AW956580 Hs.42699	EST ₈
	109737 F10078	AA055415 Hs.13233	ESTs, Moderately similar to A47582 B-cel
	110411 H48032	AW001579 Hs.9645	Homo sapiens mRNA for KIAA1741 protein,
	110660 H82117	AA782114 Hs.28043	ESTs
70	110906 N39584	AA035211 Hs.17404	ESTs
70	111018 N54067	Al287912 Hs.3628	miliogen-activated protein kinase kinase
	111091 N59858	AA300067 Hs.33032 BE301871 Hs.4867	hypothetical protein DKFZp434N185 mannosyl (alpha-1,3-)-glycoprotein beta-
	111356 N90933 111378 N93764	AW160993 Hs.326292	hypothetical gene DKFZp434A1114
	111741 R26124	AB020653 Hs.24024	KIAA0846 protein
75	111769 R27957	AW629414 Hs.24230	ESTs
-	112318 R55470	AW083384 Hs.11067	ESTs, Highly similar to T46395 hypotheti
			• • • • • • • • • • • • • • • • • • • •

	112951 T16550	AA307634 Hs.6650	vacuolar protein sorting 45B (yeast homo
	113057 T26674	AW194301 Hs.339283	Human DNA sequence from clone RP1-187J11
	113195 T57112	H83265 Hs.8881	ESTs, Weakly similar to \$41044 chromosom
_	113490 T88700	BE178110 Hs.173374	Homo sapiens cDNA FLJ10500 fis, clone NT
5	113542 T90527	H43374 Hs.7890	Homo sapiens mRNA for KIAA1671 protein,
	113803 W42789	AW880709 Hs.283683	chromosome 8 open reading frame 4
	113847 W60002	NM_005032Hs.4114	plastin 3 (T isoform)
	113910 W78175	AA113262 Hs.17901	Homo saplens, done IMAGE:3937015, mRNA,
10	113947 W84768 114047 W94427	W84768 AL035858 Hs.3807	gb:zh53d03.s1 Soares_fetal_liver_spleen_ FXYD domain-containing ion transport reg
10	115061 AA253217	AI751438 Hs.41271	Homo sapiens mRNA full length insert cDN
	115819 AA426573	AA486620 Hs.41135	endomucin-2
	115870 AA432374	NM_005985Hs.48029	snail 1 (drosophila homolog), zinc finge
	115964 AA446622	AA987568 Hs.74313	KIAA1265 protein
15	116228 AA478771	AJ767947 Hs.50841	ESTs
	116264 AA482594	D51174 Hs.272239	tysosomal
	116314 AA490588	Al799104 Hs.178705	Homo sapiens cDNA FLJ11333 fis, clone PL
	116589 D59570	Al557212 Hs.17132	ESTs, Moderately similar to 154374 gene
20	117023 H88157 117112 H94648	AW070211 Hs.102415 AW959999 Hs.293658	Homo saplens mRNA; cDNA DKFZp586N0121 (f ESTs
20	117156 H97538	W73853	ESTS
	117176 H98670	H45100 Hs.49753	uveal autoantigen with coiled coil domai
	117280 N22107	M18217 Hs.172129	Homo sapiens cDNA: FLJ21409 fis, clone C
	119559 W38197	W38197	Empirically selected from AFFX single pr
25	119866 W80814	AA496205 Hs.193700	Homo sapiens mRNA; cDNA DKFZp58610324 (f
	120655 AA287347	AA305599 Hs.238205	hypothetical protein PRO2013
	121314 AA402799	W07343 Hs.182538	phospholipid scramblase 4
	121335 AA404418 121822 AA425107	AA404418 AI743860	gb:zw37e02.s1 Soares_total_fetus_Nb2HF8_ metallothlonein 1E (functional)
30	121835 AA425435	AB033030 Hs.300670	KIAA1204 protein
-	122331 AA442872	AL133437 Hs.110771	Homo sapiens cDNA: FLJ21904 fis, clone H
	122577 AA452860	AA829725 Hs.334437	hypothetical protein MGC4248
	123160 AA488687	AA488687 Hs.284235	ESTs, Weakly similar to 138022 hypotheti
25	123486 AA599674	BE019072 Hs.334802	Homo sapiens cDNA FLJ14680 fis, clone NT
35	124059 F13673	BE387335 Hs.283713	ESTs, Weakly similar to \$64054 hypotheti
	124339 H99093 124358 N22495	H99093 Hs.343411 AW070211 Hs.102415	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep Homo sapiens mRNA; cDNA DKFZp586N0121 (f
	124364 N23031	AF265555 Hs.250646	baculoviral IAP repeat-containing 6
	124726 R15740	NM_003654Hs.104576	carbohydrate (keratan sulfate Gal-6) sul
40	124763 R39610	BE410405 Hs.76288	calpain 2, (m/li) large subunit
	125167 W45560	AL137540 Hs.102541	netrin 4
	125304 Z39833	AL359573 Hs.124940	GTP-binding protein
	125307 Z40583	AW580945 Hs.330466 AA825437 Hs.58875	ESTs ESTs
	125324 66825637		
45	125329 AA825437 125598 R66613	T40064 Hs.71968	HOIRD SAURIS HIRMA: CUNA UNFZDOGFTCOS (II
45		AA868063 Hs.104576	Homo sapiens mRNA; cDNA DKFZp564F053 (fr carbohydrate (keratan sulfate Gal-6) sul
45	125598 R66613 125609 AA868063 418245 AA128075	AA868063 Hs.104576 AA088767 Hs.83883	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced
45	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161	carbohydrate (keratan suifate Gal-6) sul transmembrane, prostate androgen induced Homo sapians cDNA FLJ13613 fis, clone PL
	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 Al051390	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 Al051390 Hs.116731	carbohydrate (keratan suifate Gal-6) sul transmembrane, prostate androgen induced Homo sapiens cDNA FLJ13613 fis, clone PL ESTs
4550	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122	AA668063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787	carbohydrate (keratan suifate Gal-6) sul transmembrane, prostate androgen induced Homo sapiens cDNA FLJ13613 fis, clone PL ESTs ESTs
	125598 R66613 125609 AA668063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1
	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602	carbohydrate (keratan suifate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr
50	125598 R66613 125609 AA668063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761	AA668063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1
	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676	AA668063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 A0051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948	carbohydrate (keratan suifate GaI-6) sul transmembrane, prostate androgen induced Homo saplens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein)
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 B2076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414	carbohydrate (keratan suifate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 12869 AA598737 128903 R69417	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076394 Hs.10509 228913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 12869 AA598737 128903 R69417 128914 AA232837	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 Al051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/nydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmaleruma vesicle associated protein
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 12869 AA598737 128903 R69417	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100862 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557	carbohydrate (keratan suifate Gal-6) sul transmembrane, prostate androgen induced Homo sapiens cDNA FL/13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shick 70kD protein 8 STAT induced STAT inhibitor 3 plasmaleruma vesicle associated protein hypothetical protein PP1057
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127556 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129087 N72695	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/nydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmaleruma vesicle associated protein
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 12869 AA598737 128903 R69417 128914 AA232837 12907 N72695 129188 M30257 129226 M96843 129255 X68277	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 228913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT Induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vasculfar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129087 N72695 129188 M30257 129226 M96843 129265 X68277 129345 AA292440	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 Al051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.10571	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT Induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascular cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible,
50	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 XU2761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129087 N72695 129188 M30257 12926 M96843 129265 X68277 129348 AA292440 129468 J03040	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 Al051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.10590 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.111779	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/nydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127556 AU051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 12893 R69417 128914 AA232837 129087 N72695 129183 M30257 129226 M96843 129255 X66277 129345 AA292440 129488 J03040 129488 AA2228107	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.100808 U82108 Hs.100818 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.10571 AW410538 Hs.111779 AW966728 Hs.54642	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascular cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, actific, cysteine-rich methionine adenosyltransferase II, beta
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AUD51390 127619 AA627122 128453 XU2761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 128903 R69417 128914 M30257 129226 M96843 129255 X66277 129345 AA292440 129488 J03040 129488 AA228107 129498 AA449789	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110577 AW410538 Hs.111577 AW366728 Hs.54642 AA449789 Hs.75511	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich methionine adenosytransferase II, beta connective tissue growth factor
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128622 L35240 128669 AA598737 128903 R69417 128914 AA232837 129937 N72695 129188 M30257 129188 M30257 12926 M96843 129265 X68277 129345 AA232440 129488 AA2328107 129488 AA2328107 129488 AA2328107 129488 AA2328107	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.117779 AW366728 Hs.54642 AA449789 Hs.75511 AL045404 Hs.46366	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich methionine adenosyltransferase II, beta connective tissue growth factor KIAA0948 protein
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AUD51390 127619 AA627122 128453 XU2761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 128903 R69417 128914 M30257 129226 M96843 129255 X66277 129345 AA292440 129488 J03040 129488 AA228107 129498 AA449789	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110577 AW410538 Hs.111577 AW366728 Hs.54642 AA449789 Hs.75511	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich methionine adenosytransferase II, beta connective tissue growth factor
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127556 A0051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128937 N72695 129183 M30257 129266 M96843 129255 X68277 129345 AA292440 129488 AA228107 129488 AA228107 129489 AA49789 12957 W1367 129619 AA610116 129627 AA258308 129762 AA450273	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.105509 Z28913 Hs.102548 W28493 Hs.105509 Z28913 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.111779 AW966728 Hs.54642 AA449789 Hs.75511 AL045404 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA453694 Hs.71968	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascular cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, actific, cysteine-rich methionine adenosyltransferase II, beta connective tissue growth factor KIAA0948 protein tetraspan NET-6 protein
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127556 AU051390 127619 AA627122 128453 XU2761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128914 AA232837 12893 R69417 128914 AA232837 129925 X68277 129226 M96843 129255 X68277 129345 AA292440 129488 AA228107 129498 AA228107 129498 AA49789 129557 W01367 129619 AA610116 129627 AA258308 129762 AA460273 129884 AA286710	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.105509 Z28913 Hs.102948 W28493 Hs.1052948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110577 AW410538 Hs.110777 AW40538 Hs.110777 AW366728 Hs.54642 AA449789 Hs.75511 AL045404 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA453694 Hs.12372 AF055581 Hs.13131	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saplens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT Induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascular cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich metholnine adenosytransferase II, beta connective tissue growth factor KIAA0948 protein tetraspan NET-6 protein Homo sapiens mRNA; cDNA DKFZp564F053 (fr triparifite motif protein TRIM2 lysosomal
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AU51390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129814 AA232837 129814 M30257 12918 M30257 12926 M96843 129265 X68277 129345 AA29240 129488 JA22840 129488 AA22840 129488 AA228107 129498 AA449789 129557 W01367 129619 AA610116 129627 AA258308 129762 AA450273 129884 AA228710 130018 T68873	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.1052948 W28493 Hs.1052948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.171695 R22497 Hs.110571 AW410538 Hs.11779 AW366728 Hs.54642 AA449789 Hs.75511 AL045404 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA209534 Hs.284243 T40064 Hs.71561 AA353093	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich methionine adenosyltransferase II, beta connective tissue growth factor KIAA0948 protein tetraspan NET-6 protein Homo saptens mRNA; cDNA DKFZp564F053 (fr tripartite motif protein TRIM2 lysosomal metallothioneln 1L
50 55 60 65	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AI051390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129814 AA232837 129815 M30257 129226 M96843 129255 X68277 129345 AA29840 129488 J03040 129488 AA228107 129498 AA249789 129557 W01367 129619 AA610116 129627 AA258308 129762 AA450273 129884 AA2286710 130018 T68873 130147 D63476	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.102948 W28493 Hs.102948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.111779 AW366728 Hs.54642 AA449789 Hs.54511 AL045404 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA209534 Hs.12372 AF055581 Hs.13131 AA353093 D63476 Hs.172813	carbohydrate (keratan sulfate GaI-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene enigma (LIM domain protein) heat shock 70kD protein 8 STAT Induced STAT Inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascular cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, actific, cysteine-rich methonine adenosyltransferase II, beta connective tissue growth factor KIAA0948 protein tetraspan NET-6 protein Homo sapiens mRNA; cDNA DKFZp564F053 (fr tripartite motif protein TRIM2 lysosomal metallothionein 1L PAK-interacting exchange factor beta
50 55 60	125598 R66613 125609 AA868063 418245 AA128075 127435 N66570 127566 AU51390 127619 AA627122 128453 X02761 128495 AF010193 128515 AA149044 128580 U82108 128623 D78676 128642 L35240 128669 AA598737 128903 R69417 128914 AA232837 129814 AA232837 129814 M30257 12918 M30257 12926 M96843 129265 X68277 129345 AA29240 129488 JA22840 129488 AA22840 129488 AA228107 129498 AA449789 129557 W01367 129619 AA610116 129627 AA258308 129762 AA450273 129884 AA228710 130018 T68873	AA868063 Hs.104576 AA088767 Hs.83883 X69086 Hs.286161 AI051390 Hs.116731 AA627122 Hs.163787 X02761 Hs.287820 NM_005904Hs.100602 BE395085 Hs.10086 U82108 Hs.101813 BE076608 Hs.105509 Z28913 Hs.1052948 W28493 Hs.1052948 W28493 Hs.180414 AW150717 Hs.345728 AW867491 Hs.107125 AI348027 Hs.108557 NM_001078Hs.109225 BE222494 Hs.180919 AA530892 Hs.171695 R22497 Hs.110571 AW410538 Hs.171695 R22497 Hs.110571 AW410538 Hs.11779 AW366728 Hs.54642 AA449789 Hs.75511 AL045404 Hs.46366 AA209534 Hs.284243 T40064 Hs.46366 AA209534 Hs.284243 T40064 Hs.71561 AA353093	carbohydrate (keratan sulfate Gal-6) sul transmembrane, prostate androgen induced Homo saptens cDNA FLJ13613 fis, clone PL ESTs ESTs fibronectin 1 MAD (mothers against decapentaplegic, Dr type I transmembrane protein Fn14 solute carrier family 9 (sodium/hydrogen CTL2 gene entgma (LIM domain protein) heat shock 70kD protein 8 STAT induced STAT inhibitor 3 plasmalemma vesicle associated protein hypothetical protein PP1057 vascutar cell adhesion molecule 1 inhibitor of DNA binding 2, dominant neg dual specificity phosphatase 1 growth arrest and DNA-damage-inducible, secreted protein, acidic, cysteine-rich methlonine adenosytransferase II, beta connective tissue growth factor KIAA0948 protein tetraspan NET-6 protein Homo saptens mRNA; cDNA DKFZp564F053 (fr tripartite motif protein TRIM2 lysosomal metallothionein 1L

	130431	L10284	AW505214	Hs.155560	calnexin
		AA243278		Hs.109059	mitochondrial ribosomal protein L12
		AA430032		Hs.252587	pituitary tumor-transforming 1
	130638	H16402	AW021276	Hs.17121	ESTs
5	130639	D59711	AJ557212	Hs.17132	ESTs, Moderately similar to 154374 gene
	130657	T94452	AW337575	Hs.201591	ESTs
		AA431571		Hs.337986	Homo saplens cDNA FLJ10934 fis, clone OV
		R79356	AF167706		cysteine-rich motor neuron 1
10		AA280375	AW190920		hypothetical protein SP329
10	130840		BE048821		small inducible cytokine subfamily A (Cy
		Z41740	AJ077288	Hs.296323	serum/glucocorticoid regulated kinase
		AA121543	AL050295		KIAA0758 protein
	131080		NM_00195		endothelin 1
15		AA101878		3Hs.303084	apelin; peptide ligand for APJ receptor
13	131091		AJ271216		dipeptidytpeptidase III
		N87590 AA256153	BE620886 AJ824144		GCN1 (general control of amino-acid synt ESTs
		W74533	AF104266		latrophilin
		U25997	NM_00315		stanniocaldin 1
20		V01512	AW939251		v-fos FBJ murine osteosarcoma viral onco
		V01512	AW939251		v-fos FBJ murine osteosarcoma viral onco
	131328	V01512	AW939251	Hs.25647	v-fos FBJ murine osteosarcoma viral onco
	131328	V01512	AW939251	Hs.25647	v-fos FBJ murine osteosarcoma viral onco
	131509	X56681	X56681	Hs.2780	jun D proto-oncogene
25	131555	AA161292	T47364	Hs.278613	interferon, alpha-inducible protein 27
	131564	AA491465	T93500	Hs.28792	Homo sapiens cDNA FLJ11041 fis, clone PL
		AA046593	AA040311	Hs.28959	ESTs
		D50914	BE559681		KIAA0124 protein
20	131756		AA443966		ESTs
30		M90657	AW960564		transmembrane 4 superfamily member 1
		W69127		3Hs.274411	SCAN domain-containing 1
		AA316186 AA384503	Al161383		ESTs, Highly similar to \$94541 1 clone 4
	_	AA136353	Al359214 Al267615	Hs.179260 Hs.38022	chromosome 14 open reading frame 4 ESTs
35		AA044755		Hs.173705	Homo sapiens cDNA: FLJ22050 fis, done H
50	132164			Hs.41270	procollagen-lysine, 2-oxoglutarate 5-dio
		AA058911	AA235709		DKFZP586O1624 protein
		AA620962		Hs.325093	Homo sapiens cDNA: FLJ21210 fis, clone C
	132314	AA285290	AF112222	Hs.323806	pinin, desmosome associated protein
40	132358	X60486	NM_003542	2Hs.46423	H4 histone family, member G
	132398	R31641	AA876616	Hs.16979	ESTs, Weakly similar to A43932 mucin 2 p
		AA489190	AW163483	Hs.48320	double ring-finger protein, Dorfin
	132490		NM_001290		LIM domain binding 2
45		AA257993	AA257992		Janus kinase 1 (a protein tyrosine kinas
45	132546		M24283	Hs.168383	intercellular adhesion molecule 1 (CD54)
	132610	AA443114	AA160511		amino acid system N transporter 2; porcu
	132840		BE379595		casein kinase 1, alpha 1 GTPase Rab14
		AA047151	BE218319 AA373314		Homo sapiens mRNA; cDNA DKFZp586P1622 (f
50	132968		AF234532		myosin X
50		AA480074	AA480074		hypothetical protein FLJ13213
	132999		Y00787	Hs.624	Interleukin 8
	133071		BE384932		ESTs, Weakly similar to AF257182 1 G-pro
	133076		AW946276		Homo sapiens mRNA; cDNA DKFZp586J021 (fr
55	133099	L09209	W16518	Hs.279518	amytoid beta (A4) precursor-like protein
	133147	D12763	AA026533	Hs.66	Interleukin 1 receptor-like 1
	133149	T16484	AA370045	Hs.6607	AXIN1 up-regulated
		AA253193	AW021103		hypothetical protein FLJ20373
C O		AA432248		Hs.183639	hypothetical protein FLJ10210
60	133220		NM_006074		Homo sapiens mRNA full length insert cDN
		AA083572	AA403045		Homo sapiens cDNA: FLJ23197 fis, clone R
	133295			Hs.213289	low density lipoprotein receptor (famili
	133349		AW631255		L-3-hydroxyacyi-Coenzyme A dehydrogenase
65	133391 133398		AW103364 NM_000499		inhibin, beta A (activin A, activin AB a cytochrome P450, subfamily I (aromatic c
05	133436		BE294068		immediate early protein
		AA090257	BE547647		hypothetical protein MGC5618
	133478			Hs.31432	cardiac ankyrin repeat protein
	133491		BE619053		eukaryotic translation Initiation factor
70		AA227913	AW880841		p53-induced protein
	133517		NM_000165		gap junction protein, alpha 1, 43kD (con
	133526		AU077051		alpha-2-macroglobulin
	133538		NM_003257		fight junction protein 1 (zona occludens
	133562			Hs.74870	H2.0 (Drosophila)-like homeo box 1
75	133584			Hs.181243	activating transcription factor 4 (tax-r
	133590	T67986	170956	Hs.75106	clusterin (complement lysis inhibitor, S

		AA148318	BE244334		ADP-ribosylation factor-like 6 Interacti
		U97105 T25747	AI301740	Hs.173381 Hs.301819	dihydropyrimidinase-like 2 zinc finger protein 146
		K02574	AW247252	113.301013	nucleoside phosphorylase
5		D78577	AI352558		tyrosine 3-monooxygenase/tryptophan 5-mo
		X53331	-	Hs.279009	matrix Gla protein
		S73591 X95735		Hs.179526	upregulated by 1,25-dihydroxyvitamin D-3
		L16862	BE410769 AW239400		zyxin G protein-coupled receptor kinase 6
10		U44975		Hs.285313	core promoter element binding protein
		M97796		Hs.180919	inhibitor of DNA binding 2, dominant neg
		U86782 AA099391	U86782	Hs.178761	26S proteasome-associated pad1 homolog
		M19267	U48959 M19267	Hs.211582 Hs.77899	myosin, light polypeptide kinase tropomyosin 1 (alpha)
15		D29992	C18356	Hs.295944	tissue factor pathway inhibitor 2
		L19314	Al125639		hairy (Drosophila)-homolog
		S78569	NM_002290		laminin, alpha 4
		U28811 L77886	NM_012201 AL034349		Golgi apparatus protein 1 protein tyrosine phosphatase, receptor t
20		C14407	AW245540		brain abundant, membrane attached signal
		M60278	AA161219		diphtheria toxin receptor (heparin-bindi
		R81509	AA102179		Homo sapiens cDNA FLJ11680 fis, done HE
		AA487558 D86962	AW580939 D86962	Hs.81875	complement component C1q receptor growth factor receptor-bound protein 10
25		AA478971	R70429	Hs.81988	disabled (Drosophila) homolog 2 (mitogen
		D50683	D50683	Hs.82028	transforming growth factor, beta recepto
		U56637	AI557280	Hs.184270	capping protein (actin filament) muscle
		M61199 M28882	AA334551 X68264	Hs.211579	sperm specific antigen 2 metanoma cell adhesion molecute
30		X15183	M30627	Hs.289088	heat shock 90kD protein 1, alpha
		S53911	NM_001773		CD34 antigen
		U20734	AU076592		jun B proto-oncogene
		D28235 AA236324	D28235 AW968058	Hs.196384 Hs 92381	prostaglandin-endoperoxide synthase 2 (p nudix (nucleoside diphosphate linked moi
35		AA148923	AL136653		decidual protein induced by progesterone
		AA174183	AK000967		KIAA1682 protein
		AA456311	AA876372		Homo sapiens mRNA; cDNA DKFZp667D095 (fr
	135071 135073	AA452000	W27190 W55956	Hs.94 Hs.94030	DnaJ (Hsp40) homolog, subfamily A, membe Homo sapiens mRNA; cDNA DKFZp586E1624 (f
40		AA282140	T53169	Hs.9587	Homo sapiens cDNA: FLJ22290 fis, clone H
	135196		C03577	Hs.9615	myosin regulatory light chain 2, smooth
		AA442054	U80983	Hs.268177	phospholipase C, gamma 1 (formerly subty
		AB000450 AB002380	AB000450 AF180681		vaccinia related kinase 2 Rho guanine exchange factor (GEF) 12
45		AB003103	AA130080		proteasome (prosome, macropain) 26S subu
		AB004884	N27852	Hs.57553	tousled-like kinase 2
		AF000573 AF008937	BE094848	Hs.15113	homogentisate 1,2-dioxygenase (homogenti
		AF009301	AF008937 AB011169	Hs 20141	syntaxin 16 similar to S. cerevisiae SSM4
50		AF009368	AF029674		KIAA1605 protein
	100113		NM_001269		chromosome condensation 1
		D00760 D11139	AA294921		v-ral simian leukemia viral oncogene hom
		D14657	AA469369 H60720	Hs.81892	tissue inhibitor of metalloproteinase 1 KIAA0101 gene product
55		D14878	AL037228		D123 gene product
		D17716	NM_002410		mannosyl (alpha-1,6-)-glycoprotein beta-
	100190 134742	D21090	M91401 NM_001346	Hs.178658	RAD23 (S. cerevisiae) homolog B diacylglycerol kinase, gamma (90kD)
	100211			Hs.123058	DEAD/H (Asp-Glu-Ala-Asp/His) box polypep
60	100238	D30742	L24959	Hs.348	catcium/calmodutin-dependent protein kin
	130283		NM_012288		TRAM-like protein
	134237 100248		D31765 NM_015156	Hs.170114	KIAA0061 protein KIAA0071 protein
	100256		D25418	Hs.393	prostaglandin (2 (prostacyclin) receptor
65	100262	D38500	D38500	Hs.278468	postmeiotic segregation increased 2-like
	134329	D38551	N92036	Hs.81848	RAD21 (S. pombe) homolog
	100281 100294		AF091035 AA331881	Hs.184627 He 75454	KIAA0118 protein peroxiredoxin 3
	100327		D55640	110.13-13-1	gb:Human monocyte PABL (pseudoautosomal
70	100335	D63391	AW247529		platelet-activating factor acetylhydrola
	134495		D63477	Hs.84087	KIAA0143 protein
	100338 135152		D86864 M96954	Hs.57735 Hs.182741	acetyl LDL receptor, SREC TIA1 cytotoxic granule-associated RNA-bi
•-	.134269	D79990	NM_014737		Ras association (RalGDS/AF-6) domain fam
75	100372		NM_014791		KIAA0175 gene product
•	134304	DOUDIU	BE613486	ms.61412	lipin 1

	100304	D84276	D84284	He ecosa	CD39 cettece (e/5)
		D86425	AW291587	Hs.66052	CD38 antigen (p45) nidogen 2
		D86978	D86978	Hs.84790	KIAA0225 protein
		D87012	D87012	Hs.194685	topolsomerase (DNA) III beta
5		D87075	AF164142		solute carrier family 23 (nucleobase tra
-		D87432	D87432	Hs.10315	solute carrier family 7 (cationic amino
		D87448	AA013051		topoisomerase (DNA) II binding protein
	134593	D87845		7Hs.234392	platelet-activating factor acetylhydrola
		HG1098-HT1098		Hs.121489	cystatin D
10	100552	HG2167-HT2237	AA019521	Hs.301946	lysosomal
		HG2415-HT2511		1Hs.231444	Homo sapiens, Similar to hypothetical pr
		HG2825-HT2949		Hs.142653	ret finger protein
		HG2887-HT3031		Hs.816	SRY (sex determining region Y)-box 2
1.5		HG4660-HT5073		Hs.103042	microtubute-associated protein 1B
15		HG4704-HT5146		Hs.172816	neuregulin 1
		HG884-HT884	AF002225	Hs.180686	ubiquitin protein ligase E3A (human papi
		HG919-HT919		Hs.166846	polymerase (DNA directed), epsilon
		J00212 J04029	J00212	M= 00036	Empirically selected from AFFX single pr
20	130149		J04029	Hs.99936 Hs.172665	keratin 10 (epidermolytic hyperkeratosis
20	131877		J04088	Hs.156346	methylenetetrahydrofolate dehydrogenase topoisomerase (DNA) II alpha (170kD)
	101016		J04543	Hs.78637	annexin A7
	134786		T29618	Hs.89640	TEK tyrosine kinase, endothelial (venous
		L07540		Hs.171075	replication factor C (activator 1) 5 (36
25	134078		L08895	Hs.78995	MADS box transcription enhancer factor 2
	101132		L11239	Hs.36993	gastrulation brain homeo box 1
		L11353	BE409525		neurofibromin 2 (bilateral acoustic neur
	332736		Z83689	Hs.114765	myeloid/lymphoid or mixed-lineage leukem
	101152	L13800	Al984625	Hs.9884	spindle pole body protein
30	135397		L14922	Hs.166563	replication factor C (activator 1) 1 (14
	432642		BE297635		heat shock 70kD protein 9B (mortalin-2)
	101168			8Hs.211569	G protein-coupled receptor kinase 5
	421155		H87879	Hs.102267	lysyl oxidase
35	101226		AF083892		tight junction protein 2 (zona occludens
33	415138		C18356	Hs.295944	tissue factor pathway inhibitor 2
	134739 130155		NM_002419	Hs.151254	mitogen-activated protein kinase kinase kallikrein 7 (chymotryptic, stratum com
	440538		W76332	Hs.79107	mitogen-activated protein kinase 14
	409916		BE313625		solute carrier family 11 (proton-coupled
40	101294		AF168418		thyroid hormone receptor interactor 4
. •	101300		BE535511		transmembrane trafficking protein
	101310		L41607	Hs.934	glucosaminyl (N-acetyl) transferase 2, I
	130344	L77566	AW250122	Hs.154879	DiGeorge syndrome critical region gene D
	101381	M13928	AW675039	Hs.1227	aminolevulinate, delta-, dehydratase
45		M13928	AW675039		aminolevulinate, delta-, dehydratase
		M14016	AW005903		uroporphyrinogen decarboxylase
		M14219	AA557660		decorin
		M15796	BE267931	Hs.78996	proliferating cell nuclear antigen
50		M21305	M21305		gb:Human alpha satellite and satellite 3
50		M22092	M22092	CU- 404C	gb:Human neural cell adhesion molecule (
		M22898 M22995	NM_000546 NM_002884		tumor protein p53 (Li-Fraumeni syndrome) RAP1A, member of RAS oncogene family
		M23379	NM_002890		RAS p21 protein activator (GTPase activa
		M24400	AW583062		chymotrypsinogen B1
55		M25753	BE280074		cyclin B1
	134116		R84694	Hs.79194	cAMP responsive element binding protein
		M28213	AA535244		RAB2, member RAS oncogene family
	130174	M29550	M29551	Hs.151531	protein phosphatase 3 (formerly 2B), cat
	129963		M29971	Hs.1384	O-6-methylguanine-DNA methyltransferase
60	132983	M30269	M30269		nidogen (enactin)
	133900		M31158	Hs.77439	protein kinase, cAMP-dependent, regulato
		M31166	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
		M31210		Hs.154210	endothelial differentiation, sphingolipi
65		M55420	S55271	Hs.247930	Epsilon , IgE
65	134691		AW382987		prostaglandin-endoperoxide synthase 1 (p
	133595			Hs.75133	transcription factor 6-like 1 (mitochond
	101700		D90337	Hs.247916	natriuretic peptide precursor C
	101714		M68874	Hs.211587	phospholipase A2, group IVA (cytosolic,
70		M74524 M80254	D28459	Hs.80612 Hs.173125	ubiquitin-conjugating enzyme E2A (RAD6 h peptidylprolyl isomerase F (cyclophilin
, ,		M81780	M80254 X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
		M81780	X59960 X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
		M81780	X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
		M81780	X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
75		M81780	X59960	Hs.77813	sphingomyelin phosphodiesterase 1, acid
	101791		M83822	Hs.62354	cell division cycle 4-like
					•

	101812	M86934	BE439894	Hs.78991	DNA segment, numerous copies, expressed	
	101813		NM_00291		replication factor C (activator 1) 2 (40	
	133396		M96326	Hs.72885	azurocidin 1 (cationic antimicrobial pro	
5	428161 129026		M96954	Hs.182741	TIA1 cytotoxic granule-associated RNA-bi	
J	101901		AL120297 H38026	Hs.308	Friend leukemia virus integration 1 arrestin 3, retinal (X-arrestin)	
	134831		AA853479		pyruvate carboxytase	
	134039		NM_00229	DHs.78672	laminin, alpha 4	
10	442355		AA456539		lysosomal-associated membrane protein 2	
10	101975 101977		AA079717 AF112213		aspartate beta-hydroxylase	
	101978		BE561610		putative Rab5-interacting protein putative transmembrane protein; homolog	
	101998		U01212	Hs.248153	olfactory marker protein	
1.5	102003		U01922	Hs.125565	translocase of inner mitochondrial membr	
15	102007		U02556	Hs.75307	t-complex-associated-testis-expressed 1-	
	102009 416658		BE245149 U03272	Hs.79432	protein tyrostne kinase 9 fibrillin 2 (congenital contractural ara	
	132951		AW821182		microfibrillar-associated protein 1	
	135389		U05237	Hs.99872	fetal Alzheimer antigen	
20	102048		U07225	Hs.339	purinergic receptor P2Y, G-protein coupl	
	130145		U34820	Hs.151051	mitogen-activated protein kinase 10	
	303153 420269		U09759 U72937	Hs.246857 Hs.96264	mitogen-activated protein kinase 9 alpha thalassemla/mental retardation syn	
	102095		U11313	Hs.75760	sterol carrier protein 2	
25	102123		NM_00180		centromere protein A (17kD)	
	102126		AW950870		protein phosphalase 1, regulatory (inhib	
	102133 102139		AU076845		BCL2/adenovirus E1B 19kD-interacting pro	
	102162		NM_004419 AA450274		dual specificity phosphatase 5 CDC16 (cell division cycle 16, S. cerevi	
30	102164		NM_000107		damage-specific DNA binding protein 2 (4	
	427653		AA159001	Hs.180069	nuclear respiratory factor 1	
	131817		U20536	Hs.3280	caspase 6, apoptosis-related cysteine pr	
	102200 102210		AA232362 BE619413		branched chain aminotransferase 1, cytos eukaryotic translation initiation factor	
35	102214		U23752	Hs.32964	SRY (sex determining region Y)-box 11	
	132811		U25435	Hs.57419	CCCTC-binding factor (zinc finger protei	
	131319		NM_00315		stanniocalcin 1	
	102256 132316		U28251	Hs.53237	ESTs, Highly similar to Z169_HUMAN ZINC	
40	102269		U28831 U30245	Hs.44566	KIAA1641 protein gb:Human myelomonocytic specific protein	
	417526		AA568906	Hs.82240	syntaxin 3A	
	102293		AF090116		regulator of G-protein signalling 7	
	102298		AA382169		N-myc (and STAT) interactor	
45	102325 (428734 (AI815867 BE303044	Hs.50130 Hs 192023	necdin (mouse) homolog eukaryotic translation initiation factor	
	102361		AA223616		chromosome 11 open reading frame 4	
	102367		U39656	Hs.118825	mitogen-activated protein kinase kinase	
	102388		AA362907		proline arginine-rich end leucine-rich r	
50	102394 129829		NM_003816 AF010258		a disintegrin and metalloproteinase doma homeo box A9	
50	102409		BE300330		selenophosphate synthetase 2	
	133746		AW410035		MAD (mothers against decapentaplegic, Dr	
	102423		Z47542	Hs.179312	small nuclear RNA activating complex, po	
55	132828		AB014615		fibroblast growth factor 8 (androgen-ind	
55	132828 I		AB014615 AB014615		fibroblast growth factor 8 (androgen-ind fibroblast growth factor 8 (androgen-ind	
	132828		AB014615		fibroblast growth factor 8 (androgen-ind	
	425322		U63630	Hs.155637	protein kinase, DNA-activated, catalytic	
60	102450		U48251	Hs.75871	protein kinase C binding protein 1	
60	129350 I 102534 I		U50535 U96759	Hs.110630 Hs.198307	Human BRCA2 region, mRNA sequence CG006 von Hippel-Lindau binding protein 1	
	130457		AB014595		aullin 48	
	135065		AA019401		cyclic nucleotide gated channel beta 1	
<i></i>	102560			Hs.63984	cadherin 13, H-cadherin (heart)	
65	102567			Hs.146847	TRAF family member-associated NFKB activ	
	417173 \ 102638 \		U61397 U67319	Hs.81424 Hs.9216	ubiquitin-like 1 (sentrin) caspase 7, apoptosis-related cysteine pr	
	132736		AW081883		Homo sapiens cDNA: FLJ23037 fis, clone L	
	133070		U92649	Hs.64311	a disintegrin and metalloproteinase doma	
70	102663		NM_002270	Hs.168075	karyopherin (importin) beta 2	
	134660 (U73524	Hs.87465	ATP/GTP-binding protein	
	102735 U 102741 U		AF111106 AW959829		protein phosphatase 4, regulatory subuni hypothetical protein MGC14433	
	130564		U82671	Hs.36980	melanoma antigen, family A, 2	
75	130564	U82671	U82671	Hs.36980	melanoma antigen, family A, 2	
	132164 (AI752235	Hs.41270	procollagen-tysine, 2-oxogtutarate 5-dio	
	•	•				

	102823 U90914	D85390 Hs.5057	carboxypeptidase D
	102826 U91316	NM_007274Hs.8679	cytosolic acyl coenzyme A thioester hydr
	102831 U91932	AA262170 Hs.80917	adaptor-related protein complex 3, sigma
_	102846 U96131	BE264974 Hs.6566	thyroid hormone receptor interactor 13
5	129777 U97018	U97018 Hs.12451	echinoderm microtubule-associated protei
	134161 U97188	AA634543 Hs.79440	IGF-II mRNA-binding protein 3
	134854 V00503	J03464 Hs.179573	collagen, type I, alpha 2
	429257 X04327	AW163799 Hs.198365	2,3-bisphosphoglycerate mutase
10	413985 X06389 419768 X07496	Al018666 Hs.75667 T72104 Hs.93194	synaptophysin
10	102915 X07820	X07820 Hs.2258	apolipoprotein A-I matrix metalloproteinase 10 (stromelysin
	134656 X14787	Al750878 Hs.87409	thrombospondin 1
	413858 X15525	NM_001610Hs.75589	acid phosphatase 2, lysosomal
	102968 X16396	AU076611 Hs.154672	methylene tetrahydrofolate dehydrogenase
15	102971 X16609	X16609 Hs.183805	ankyrin 1, erythrocytic
	134037 X53586	Al808780 Hs.227730	integrin, alpha 6
	134037 X53586	Al808780 Hs.227730	integrin, alpha 6
	103023 X53793	AW500470 Hs.117950	multifunctional polypeptide similar to S
20	103037 X54936 130282 X55740	BE018302 Hs.2894 BE245380 Hs.153952	placental growth factor, vascular endoth 5' nucleotidase (CD73)
20	134542 X57025	M14156 Hs.85112	insulin-like growth factor 1 (somatomedi
	128568 X60673	H12912 Hs.274691	adenylate kinase 3
	128568 X60673	H12912 Hs.274691	adenylate kinase 3
	103093 X60708	S79876 Hs.44926	dipeptidylpeptidase IV (CD26, adenosine
25	413076 X62048	U10564 Hs.75188	wee1 (S. pombe) homolog
	129063 X63097	X63094 Hs.283822	Rhesus blood group, D antigen
	424460 X63563	BE275979 Hs.296014	polymerase (RNA) II (DNA directed) polyp
	411077 X64037	AW977263 Hs.68257	general transcription factor IIF, polype
30	103181 X69636 103184 X69878	X69636 Hs.334731 U43143 Hs.74049	Horno saplens, clone IMAGE:3448306, mRNA, fms-related tyrosine kinase 4
50	103194 X70649	NM_004939Hs.78580	DEAD/H (Asp-Giu-Ala-Asp/His) box polypep
	103208 X72841	AW411340 Hs.31314	retinoblastoma-binding protein 7
	129698 X74987	BE242144 Hs.12013	ATP-binding cassette, sub-family E (OABP
2.5	131486 X83107	F06972 Hs.27372	BMX non-receptor tyrosine kinase
35	130729 X84194	Al963747 Hs.18573	acytphosphatase 1, erythrocyte (common)
	103334 X85753 132645 X87870	NM_001260Hs.25283	cyclin-dependent kinase 8
	135094 X89066	Al654712 Hs.54424 NM_003304Hs.250687	hepatocyte nuclear factor 4, alpha transient receptor potential channel 1
	103352 X89398	H09366 Hs.78853	uracii-ONA glycosylase
40	103352 X89398	H09366 Hs.78853	uracii-DNA glycosylase
	103353 X89399	X89399 Hs.119274	RAS p21 protein activator (GTPase activa
	132173 X89426	X89426 Hs.41716	endothelial cell-specific molecule 1
	103371 X91247 131584 X91648	X91247 Hs.13046 AA598509 Hs.29117	thioredoxin reductase 1 purine-rich element binding protein A
45	103376 X92098	AL036166 Hs.323378	coated vesicle membrane protein
	103378 X92110	AL119690 Hs.153618	HCGVIII-1 protein
	128510 X94703	X94703	RAB28, member RAS oncogene family
	103410 X96506	AA158294 Hs.295362	DR1-associated protein 1 (negative cofac
60	133490 X97230	AF022044 Hs.274601	killer cell immunoglobulin-like receptor
50	332689 X97230	AF022044 Hs.274601	killer cell immunoglobulin-like receptor
	103438 X98263 103440 X98296	AW175781 Hs.152720 X98296 Hs.77578	M-phase phosphoprotein 6
	103452 X99584	X98296 Hs.77578 NM_006936Hs.85119	ubiquitin specific protease 9, X chromos SMT3 (suppressor of mif two 3, yeast) ho
	133536 Y00264	W25797.comp	Hs.177486 amyloid beta (A4) precursor protein (pro
55	420234 Y07566	AW404908 Hs.96038	Ric (Drosophila)-like, expressed in many
	426502 Y07759	Y07759 Hs.170157	myosin VA (heavy polypeptide 12, myoxin)
	134662 Y07827	NM_007048Hs.284283	butyrophilin, subfamily 3, member A1
•	132083 Y07867	BE386490 Hs.279663	Pirin
60	103500 Y09443 134389 Y09858	AW408009 Hs.22580	alkylglycerone phosphate synthase
oo	132084 Y12394	Y09858 Hs.82577 NM_002267Hs.3886	spindlin-like karyopherin alpha 3 (importin alpha 4)
	103540 Z11559	NM_002197Hs.154721	aconitase 1, soluble
	133152 Z11695	Z11695 Hs.324473	mitogen-activated protein kinase 1
	103548 Z15005	Z15005 Hs.75573	centromere protein E (312kD)
65	103612 Z46261	BE336654 Hs.70937	H3 histone family, member A
	129092 AA011243	D56365 Hs.63525	poly(rC)-binding protein 2
	103692 AA018418	AW137912 Hs.227583	Homo sapiens chromosome X map Xp11.23 L-
	103695 AA018758 129796 AA018804	AW207152 Hs.186600 BE218319 Hs.5807	ESTs GTPase Rab14
70	434993 AA031993	AA306325 Hs.4311	SUMO-1 activating enzyme subunit 2
. 5	132683 AA044217	BE264633 Hs.143638	WD repeat domain 4
	131887 AA046548	W17064 Hs.332848	SWI/SNF related, matrix associated, acti
	103723 AA057447	BE274312 Hs.214783	Homo sapiens cDNA FLJ14041 fis, clone HE
75	453368 AA058376	W20296 Hs.288178	Homo sapiens cDNA FLJ11968 fis, clone HE
75	133260 AA083572	AA403045 Hs.6906	Homo saplens cDNA: FLJ23197 fis, clone R
	103765 AA085696	AA085696 Hs.169600	KIAA0826 protein

	400700 4		41000000	11: 404405	NO.
	103766 A			Hs.191435	ESTs
	103767 A		BE244667	400445	CGI-100 protein
	132051 A		AA393968		HSPC030 protein
5	103773 A			Hs.101077	ESTs, Wealty similar to T22363 hypotheti
3	135289 A		AW372569		hypothetical protein MGC10924 similar to
	409659 A		AW970843		eukaryotic translation initiation factor
	103794 A		AF244135		hepatocellular carcinoma-associated anti
	131471 A		AA164842		KIAA1600 protein
10	134319 A		BE304999		fumarate hydratase
10	103807 A		AW958264		similar to yeast Upf3, variant B
	446392 A		AF142419		homolog of mouse quaking QKI (KH domain
	129863 A		BE379765		sperm associated antigen 9
	103850 A		AA187101	MS.213194	hypothetical protein MGC10895
15	103855 A		W02363	11- 4044	hypothetical protein FLJ10330
15	103861 A		AA206236		hypothetical protein FLJ12783
	130634 A			Hs.127824	ESTs, Weakly similar to T28770 hypotheti
	447735 A		AA775268		Homo saplens cDNA: FLJ23020 fis, done L
	103909 A		AA249611		SH3 domain binding glutamic acid-rich pr
20	458928 A/		AF043117		ubiquitination factor E4B (homologous to
20	415824 A/		D42039	Hs.78871	mesoderm development candidate 2
	129013 A		AA371156		DKFZP564M112 protein
	129435 A		AF151852		CGI-94 protein
	103988 A		AA314389		ADP-ribosylation factor-like 5
25	104000 A			Hs.80475	polymerase (RNA) II (DNA directed) polyp
25	425284 A/		AF155568		NS1-associated protein 1
	128629 A/		AL096748		DKFZP434A043 protein
	133281 A/		AK001601		high-mobility group 20A
	104104 A		AA422029		ESTs, Wealty similar to hyperpolarizatio
30	332455 A/		NM_005754	NS.220009	Ras-GTPase-activating protein SH3-domain
50	132091 A/ 135073 A/		AW954243	Un 04020	KIAA0251 protein
				Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
	131367 A/ 129593 A/			Hs.173933	nuclear factor I/A
	133505 C			Hs.98314 Hs.324504	Homo sapiens mRNA; cDNA DKFZp586L0120 (f Homo sapiens mRNA; cDNA DKFZp586J0720 (f
35	132064 C		Al630124 AA121098		serum-inducible kinase
55	442351 C			Hs.8261	hypothetical protein FLJ22393
	131427 CI		AF151879		CGI-121 protein
	433892 C	-		Hs.323966	Homo sapiens clone H63 unknown mRNA
	104282 C			Hs.332338	EST
40	134827 D1		BE314037		coproporphyrinogen oxidase (coproporphyr
-10	425330 D2			Hs.155650	KIAA0014 gene product
	131742 D		AA961420		ESTs
	456935 D		AA370362		EGF-TM7-latrophilin-related protein
	425218 D8		NM_014909		KIAA1036 protein
45	104334 D			Hs.78771	phosphoglycerate kinase 1
	134593 D8		NM_000437		platelet-activating factor acetylhydrola
	134731 D8			Hs.89404	msh (Drosophila) homeo box homolog 2
	445776 H		NM 001310		cAMP responsive element binding protein-
	131670 H		_	Hs.15589	ESTs
50	104394 H		AA129551		Homo saplens cDNA: FLJ21409 fis, clone C
	104402 H			Hs.132956	ESTs
	439130 H7			Hs.124707	ESTs
	129077 H7			Hs.108479	ESTs
	104417 H			Hs.320861	Kruppel-like factor 8
55	134927 L3			Hs.91296	Integrin, alpha 8
	129280 M	63154	M63154	Hs.110014	gastric intrinsic factor (vitamin B synt
	134498 M	63180	AW246273	Hs.84131	threonyl-tRNA synthetase
	104460 M	91504	AW955705	Hs.62604	Homo sapiens, clone IMAGE:4299322, mRNA,
	104488 NS	56191	N56191	Hs.106511	protocadherin 17
60	131248 N7	78483	A1038989	Hs.332633	Bardet-Biedl syndrome 2
	130017 R1	14652	AK000096	Hs.143198	inhibitor of growth family, member 3
	104530 R2	20459	AK001676	Hs.12457	hypothetical protein FLJ10814
	104534 R2	22303	R22303		gb:yh26b09.r1 Soares placenta Nb2HP Homo
	104544 R3	33779	Al091173	Hs.222362	ESTs, Weakly similar to p40 [H.sapiens]
65	133328 R3	36553	AW452738	Hs.265327	hypothetical protein DKFZp761I141
	104567 R6	64534	AA040620		hypothetical protein AF140225
	129575 R7			Hs.278428	progestin induced protein
	130776 R7		AF167706	Hs.19280	cysteine-rich motor neuron 1
a ^	104599 R8		AW815036		ESTs
70	104660 AA		BE298665		Homo sapiens mRNA; cDNA DKFZp564D016 (fr
	104667 AA		Al239923		ESTs
	104718 A	A018409	Al143020		ESTs, Weakly similar to 138022 hypotheti
	104764 AA			Hs.278585	ESTs
	104786 AA		AA027167	Hs.10031	KIAA0955 protein
75	104787 A		AA027317		gb:ze97d11.s1 Soares_fetal_heart_NbHH19W
	134079 A/	A029423	AK001751	Hs.171835	hypothetical protein FLJ 10889

	404004 44004057	A1050700 U- 04000	DOT: W. da.d B. d MANAGERIA
	104804 AA031357	AI858702 Hs.31803	ESTs, Wealty similar to N-WASP [H.sapten
	104865 AA045136	T79340 Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
	130828 AA053400	AW631469 Hs.203213	ESTS
5	104907 AA055829	AA055829 Hs.196701	ESTs, Weakly similar to ALU1_HUMAN ALU S
,	104943 AA065217	AF072873 Hs.114218 H63789 Hs.296288	frizzled (Drosophila) homolog 6
	105013 AA116054 105024 AA126311	AA126311 Hs.9879	ESTs, Weakly similar to KIAA0638 protein
	132592 AA129390	AW803564 Hs.288850	ESTS
	105038 AA130273	AW503733 Hs.9414	Homo sapiens cDNA: FLJ22528 fis, clone H
10	105077 AA142919	W55946 Hs.234863	KIAA1488 protein
10	105096 AA150205	AL042506 Hs.21599	Homo sapiens cDNA FLJ12082 fis, clone HE
	129215 AA176867	AB040930 Hs.126085	Kruppel-like factor 7 (ubiquitous)
	105169 AA180321	BE245294 Hs.180789	KIAA1497 protein S164 protein
		NM 006283Hs.173159	
15	132796 AA180487 427210 AA187634	BE396283 Hs.173987	transforming, acidic colled-coll contain
13	105200 AA195399	AA328102 Hs.24641	eukaryotic translation initiation factor cytoskeleton associated protein 2
	130114 AA234717	AA233393 Hs.14992	
	105330 AA234743	AW338625 Hs.22120	hypothetical protein FLJ11151 ESTs
	105337 AA234957	A1468789 Hs.347187	myotubularin related protein 1
20	422040 AA235604	AA172106 Hs.110950	Rag C protein
20	105376 AA236559	AW994032 Hs.8768	hypothetical protein FLJ10849
	105397 AA242868	AA814807 Hs.7395	hypothetical protein FLJ23182
	431679 AA251776	AK000046 Hs.343877	hypothetical protein FLJ20039
	131991 AA251909	AF053306 Hs.36708	budding uninhibited by benzimidazoles 1
25	421305 AA252672	BE397354 Hs.324830	diptheria toxin resistance protein requi
25	105489 AA256157	AA256157 Hs.24115	Homo sapiens cONA FLJ14178 fis, clone NT
	105508 AA256680	AA173942 Hs.326416	Homo saplens mRNA; cDNA DKFZp564H1916 (f
	105539 AA258873	AB040884 Hs.109694	KIAA1451 protein
	135172 AA262727	AB028956 Hs.12144	KIAA1033 protein
30	131569 AA281451	AL389951 Hs.271623	nucleoporin 50kD
-	431129 AA281545	AL137751 Hs.263671	Homo sapiens mRNA; cDNA DKFZp434l0812 (f
	105643 AA282069	BE621719 Hs.173802	KIAA0603 gene product
	105659 AA283044	AA283044 Hs.25625	hypothetical protein FLJ11323
	105666 AA283930	AA426234 Hs.34906	ESTs, Weakly similar to T17210 hypotheti
35	105674 AA284755	AI609530 Hs.279789	histone deacetylase 3
	105709 AA291268	Al928962 Hs.26761	DKFZP586L0724 protein
	105722 AA291927	Al922821 Hs.32433	ESTs
	105765 AA343514	AA299688 Hs.24183	ESTs
	115951 AA398109	BE546245 Hs.301048	sec13-like protein
40	130884 AA398109	BE546245 Hs.301048	sec13-like protein
	105962 AA405737	AW880358 Hs.339808	hypothetical protein FLJ10120 .
	105985 AA406610	AA406610	gb:zv15b10.s1 Soares_NhHMPu_S1 Homo sapi
	106008 AA411465	AB033888 Hs.8619	SRY (sex determining region Y)-box 18
4.5	457322 AA416886	Al815486 Hs.243901	Homo sapiens cDNA FLJ20738 fis, done HE
45	134222 AA424013	AW855861 Hs.8025	Homo sapiens clone 23767 and 23782 mRNA
	446954 AA424148	AB037850 Hs.16621	DKFZP434l116 protein
	106141 AA424558	AF031463 Hs.9302	phosducin-like
	447973 AA424961	AB011169 Hs.20141	similar to S. cerevisiae SSM4
50	106157 AA425367	W37943 Hs.34892	KIAA1323 protein
50	428314 AA425921	AW135049 Hs.26285	Homo sapiens cDNA FLJ10643 fis, clone NT
	446727 AA426220	AB011095 Hs.16032	KIAA0523 protein
	106196 AA427735	AA525993 Hs.173699	ESTs, Weakly similar to ALU1_HUMAN ALU S
	457714 AA430673	AA083764	hypothetical protein MGC3178
EE	133200 AA432248	AB037715 Hs.183639	hypothetical protein FLJ10210
55	106302 AA435896	AA398859 Hs.18397	hypothetical protein FLJ23221
	106328 AA436705	AL079559 Hs.28020	KIAA0766 gene product
	450534 AA446561	Al570189 Hs.25132	KIAA0470 gene product
	106423 AA448238	AB020722 Hs.16714	Rho guanine exchange factor (GEF) 15
60	439608 AA449756 106477 AA450303	AW864696 Hs.301732 R23324 Hs.41693	hypothetical protein MGC5306 DnaJ (Hsp40) homolog, subfamily B, membe
00	106503 AA452411	R23324 Hs.41693 AB033042 Hs.29679	cofactor required for Sp1 transcriptiona
	446999 AA454566	AA151520	
	106543 AA454667	AA676939 Hs.69285	hypothetical protein MGC4485 neuroptlin 1
	442007 AA456437	AA301116 Hs.142838	nucleolar phosphoprotein Nopp34
65	106589 AA456646	AK000933 Hs.28661	Homo sapiens cDNA FLJ10071 fis, clone HE
05	106593 AA456826	AW296451 Hs.24605	ESTs
	106596 AA456981	AA452379	ESTs, Moderately similar to ALU7_HUMAN A
	423064 AA458959	AF265208 Hs.8740	SWI/SNF related, matrix associated, acti
	106636 AA459950	AW958037 Hs.286	ribosomal protein L4
70	106654 AA460449	AW075485 Hs.286049	phosphoserine aminotransferase
, ,	131353 AA463910	AW754182	gb:RC2-CT0321-131199-011-c01 CT0321 Homo
	106707 AA464603	AK000566 Hs.98135	hypothetical protein FLJ20559
	452909 AA464606	NM_015368Hs.30985	pannexin 1
	106717 AA465093	AA600357 Hs.239489	TIA1 cytotoxic granule-associated RNA-bi
75		AB014548 Hs.31921	KIAA0648 protein
1.3	453141 AA465692	WD0 14040 D221251	VIVADAR bioresi
15	453141 AA465692 106747 AA476473	NM_007118Hs.171957	triple functional domain (PTPRF interact

	106773 AA478109	AA478109 Hs.188833	ESTs ·
	106781 AA478474	AA330310 Hs.24181	ESTs
	106817 AA480889	D61216 Hs.18672	ESTs
	106846 AA485223	AB037744 Hs.34892	KIAA1323 protein
5	106848 AA485254	AA449014 Hs.121025	chromosome 11 open reading frame 5
	106856 AA486183	W58353 Hs.285123	Homo sapiens mRNA full length insert cDN
	418699 AA496936	BE539639 Hs.173030	ESTs, Weakly similar to ALU8_HUMAN ALU S
	107001 AA598589	Al926520 Hs.31016	putative DNA binding protein
10	442853 AA598831	AW021276 Hs.17121	ESTs
10	107054 AA600150	Al076459 Hs.15978	KIAA1272 protein
	107059 AA608545 107080 AA609210	BE614410 Hs.23044 AL122043 Hs.19221	RAD51 (S. cerevislae) homolog (E coli Re hypothetical protein DKFZp566G1424
	107115 AA610108	BE379623 Hs.27693	peptidylprotyl isomerase (cyclophilin)-l
	107130 AA620582	AB033106 Hs.12913	KIAA1280 protein
15	107156 AA621239	AA137043 Hs.9663	programmed cell death 6-Interacting prot
	107174 AA621714	BE122762 Hs.25338	ESTs
	130621 AA621718	AW513087 Hs.16803	LUC7 (S. cerevislae)-like
	107190 D19673	AA836401 Hs.87860	ESTs
20	132626 D25755	AW504732 Hs.21275	hypothetical protein FLJ11011
20	107217 D51095	AL080235 Hs.35861	DKFZP586E1621 protein
	332584 D60272 444655 T08879	AA357879 Hs.29423 AF088886 Hs.11590	ESTs; Weakly similar to macrophage lecti
	107295 T34527	AA186629 Hs.80120	cathepsin F UDP-N-acetyl-alpha-D-galactosamine:polyp
	107299 T40327	BE277457 Hs.30661	hypothetical protein MGC4606
25	107315 T62771	AA316241 Hs.90691	nudeophosmin/nudeoplasmin 3
	107316 T63174	T63174 Hs.193700	Homo sapiens mRNA; cDNA DKFZp586I0324 (f
	107328 T83444	AW959891 Hs.76591	KIAA0887 protein
	107334 T93641	T93597 Hs.187429	ESTs
20	456340 U48263	U48263 Hs.89040	prepronociceptin
30	128636 U49065	U49065 Hs.102865	interleukin 1 receptor-like 2
	129938 U79300 107375 U88573	AW003668 Hs.135587 BE011845 Hs.251064	Human clone 23629 mRNA sequence high-mobility group (nonhistone chromoso
	130074 U93867	AL038596 Hs.250745	polymerase (RNA) III (DNA directed) (62k
	107387 W01094	D86983 Hs.118893	Melanoma associated gene
35	132036 W01568	AL157433 Hs.37706	hypothetical protein DKFZp434E2220
	107426 W26853	W26853 Hs.291003	hypothetical protein MGC4707
	135388 W27965	W27965 Hs.99865	epimorphin
	130419 W36280	AF037448 Hs.155489	NS1-associated protein 1
40	107469 W47063 434203 W79060	W47063 Hs.94668 BE262677 Hs.283558	ESTs hypothetical protein PRO1855
+0	107506 W88550	AB028981 Hs.8021	KIAA1058 protein
	132358 X60486	NM_003542Hs.46423	H4 histone family, member G
	107522 X78931	X78931 Hs.99971	zinc finger protein 272
4.5	456495 Z14077	NM_003403Hs.97496	YY1 transcription factor
45	107582 AA002147	AA002147 Hs.59952	EST
	107609 AA004711	R75654 Hs.164797	hypothetical protein FLJ13693
	107661 AA010383	AA010383 Hs.60389 AA015761 Hs.60642	ESTs ESTs
	107714 AA015761 107775 AA018772	AW008846 Hs.60857	ESTs
50	107832 AA021473	AA021473	gb:ze66c11.s1 Soares retina N2b4HR Homo
	107859 AA024835	AW732573 Hs.47584	potassium voltage-gated channel, delayed
	107914 AA027229	AA027229 Hs.61329	ESTs, Weakly similar to T16370 hypotheti
	107935 AA029428	AA029428 Hs.61555	ESTs
55	410196 AA035143	Al936442 Hs.59838	hypothetical protein FLJ10808
22	131461 AA035237 108007 AA039347	AA992841 Hs.27263 AA039347 Hs.61916	KIAA1458 protein EST
	108029 AA040740	AA040740 Hs.62007	ESTs
	108040 AA041551	AL121031 Hs.159971	SWI/SNF related, matrix associated, acti
	108084 AA045513	AA058944 Hs.116602	Homo sapiens, done IMAGE:4154008, mRNA,
60	108088 AA045745	AA045745 Hs.62886	ESTs
	108168 AA055348	Al453137 Hs.63176	ESTs
	130719 AA056582	AA679262 Hs.14235	hypothetical protein FLJ20008; KIAA1839
	108189 AA056697 108190 AA056746	AW376061 Hs.63335 AA056746 Hs.63338	ESTs, Moderately similar to A46010 X-lin EST
65	108203 AA057678	AW847814 Hs.289005	Homo sapiens cDNA: FLJ21532 fis, done C
05	108216 AA058681	AA524743 Hs.44883	ESTs
	108217 AA058686	AA058686 Hs.62588	ESTs
	108245 AA062840	BE410285 Hs.89545	proteasome (prosome, macropain) subunit,
70	108277 AA064859	AA064859	gb:zm50f03.s1 Stratagene fibroblast (937
70	108280 AA065069	AA065069	gb:zm12e11.s1 Stratagene pancreas (93720
	108309 AA069923	AA069818 AA069820 Hs.180909	gb:zm67e03.r1 Stratagene neuroepithelium peroxiredoxin 1
	108340 AA070815 108403 AA075374	AA075374	gb:zm87a01.s1 Stratagene ovarian cancer
	108427 AA076382	AA076382	gb:zm91g08.s1 Stratagene ovarian cancer
75	108435 AA078787	T82427 Hs.194101	Homo sapiens cDNA: FLJ20869 fis, clone A
	108439 AA078986	AA078986	gb:zm92h01.s1 Stratagene ovarian cancer

	109466	AA079393	AA079393	Un 2400	and advances a coldess code with MI-
		AA079393 AA079487	AA079487	⊓S.340Z	cytochrome c oxidase subunit VIIc
		AA083207		Hs.68270	gb:zm97f08.s1 Stratagene colon HT29 (937 EST
		AA083256	AA083256	113.50210	gb:zn08g12.s1 Stratagene hNT neuron (937
5		AA084415	AA084415		gb:zn06g09.s1 Stratagene hNT neuron (937
_		AA085274	AA100796		gb:zm26c06.s1 Stratagene pancreas (93720
		AA088678	AI732404	Hs.68846	ESTs
-		AA100925	AJ907537	Hs.76698	stress-associated endoplasmic reticulum
	432645	AA101255	D14041	Hs.347340	H-2K binding factor-2
10	130385	AA126474	AW067800	Hs.155223	stanniocalcin 2
	108749	AA127017	AA127017	Hs.71052	ESTs
		AA129968	Al652236	Hs.49376	hypothetical protein FLJ20644
		AA130240	AA045088	Hs.62738	ESTs
		AA131866	AF188527		ESTs, Weakly similar to AF174605 1 F-box
15		AA132983	AL117452		DKFZP586G1517 protein
		AA133250	AK001468		anillin (Drosophila Scraps homolog), act
		AA133583	L46353	Hs.2726	high-mobility group (nonhistone chromoso
		AA135941	AK001431	MS.5105	hypothetical protein FLJ10569
20		AA148650	AA148650	LI- 100000	gb:zo09e06.s1 Stratagene neuroepithelium
20		AA151110 AA155754	A1304870	Hs.188680 Hs.332436	ESTs EST
		AA156125	A1056548		
		AA156289	Al611807		hypothetical protein FLJ20992 similar to hypothetical protein FLJ13397
		AA156997	AA156755		ESTs
25		AA157291	AA157291		ubinudein 1
		AA157293	AA157293		ESTs
		AA164293	AA164293		ESTs
	109072	AA164676	AI732585	Hs.22394	hypothetical protein FLJ10893
	426981	AA167375	AL044675	Hs.173081	KIAA0530 protein
30	130346	AA167550	H05769	Hs.188757	Homo saplens, clone MGC:5564, mRNA, comp
	109146	AA176589		Hs.142078	EST
		AA180448		Hs.144300	EST
		AA187144	NM_00195		endothelin 1
25		AA189170		Hs.109441	MSTP033 protein
35		AA192757		Hs.333512	similar to rat myomegalin
		AA205650 AA233342		Hs.170142 Hs.289069	ESTs
		AA233472	BE619092		hypothetical protein FLJ21016
		AA233472 AA234110	AI471639		Homo sapiens cDNA: FLJ21869 fis, clone H ESTs
40	109537		AI858695	Hs.34898	ESTs
	109556		AI925294	Hs.87385	ESTs
	109577		F02206	Hs.296639	Homo sapiens potassium channel subunit (
	109578		F02208	Hs.27214	ESTs
	109595	F02544	AA078629	Hs.27301	ESTs
45	109625	F03918	H29490	Hs.22697	ESTs
	428376		AF119665	Hs.184011	pyrophosphatase (inorganic)
•	109648		H17800	Hs.7154	ESTs
	109671		R59210	Hs.26634	ESTs
50	109699		H18013	Hs.167483	ESTs
50	109820			Hs.119021	ESTs
	109933		R52417	Hs.20945	Homo sapiens clone 24993 mRNA sequence
	110014 110039			Hs.7242	Homo saptens mRNA full length insert cDN
•	110039		H11938 R44557	Hs.21907 Hs.23748	histone acetyltransferase ESTs
55	110107		AW151660		ESTs
55	110155		Al559626	Hs.93522	Homo saptens mRNA for KIAA1647 protein,
	110197		AW090386		arrestin, beta 1
	110223		H19836	Hs.31697	ESTs
	110306		H38087	Hs.105509	CTL2 gene
60	110335		H65490	Hs.18845	ESTs .
	110342	H40567	H40961	Hs.33008	ESTs
	110395	H46966	AA025116	Hs.33333	ESTs
	110511		H56640	Hs.221460	ESTs
~	110523		A1040384	Hs.19102	ESTs, Weakly similar to organic anion tr
65	110715		H96712	Hs.269029	ESTs
	110754		AW302200		KIAA0672 gene product
	428454		U55936	Hs.184376	synaptosomal-associated protein, 23kD
	431663		NM_016569		TBX3-iso protein
70	134263		AW973443		RNA (guanine-7-) methyttransferase
70	110938		N48982	Hs.38034	Homo sapiens cDNA FLJ12924 fis, clone NT
	110983 111081		NM_015367		MIL1 protein
	111128		Al146349 AW505364		CGI-112 protein LATS (targe tumor suppressor, Drosophila
	431548		AI834273		novel protein
75	111216		AW139408		ESTs
	437562		AB001636		DEAD/H (Asp-Ghi-Ala-Asp/His) box polypep
					t new houtheh

	444200 000420	AMP/70778 Un 40067	ECT-
	111399 R00138 111514 R07998	AW270776 Hs.18857 R07998	ESTs
	428744 R08929	BE267033 Hs.192853	gb:yf16g11.s1 Soares fetal liver spleen ublquttin-conjugating enzyme E2G 2 (homo
	111574 R10307	AI024145 Hs.188526	
5	111804 R33354	AA482478 Hs.181785	ESTs ESTs
,	111831 R36083	R36095 Hs.268695	ESTs
	426773 R37938	NM_015556Hs.172180	KIAA0440 protein
	111904 R39330	Z41572	gb:HSCZYB122 normalized Infant brain cDN
	428371 R40816	AB012193 Hs.183874	cullin 4A
10	112033 R43162	R49031 Hs.22627	ESTs
- •	130987 R45698	BE613269 Hs.21893	hypothetical protein DKFZp761N0624
	112300 R54554	H24334 Hs.26125	ESTs
	112513 R68425	R68425 Hs.13809	hypothetical protein FLJ10648
	112514 R68568	R68568 Hs.183373	src homology 3 domain-containing protein
15	112522 R68763	R68857 Hs.265499	ESTs
	112540 R70467	R69751	gb:yi40a10.s1 Soares placenta Nb2HP Homo
	428655 R73565	H05769 Hs.188757	Homo sapiens, clone MGC:5564, mRNA, comp
	129534 R73640	AK002126 Hs.11260	hypothetical protein FLJ11264
	112597 R78376	R78376 Hs.29733	EST
20	112732 R92453	R92453 Hs.34590	ESTs
	451798 T03865	BE297567 Hs.27047	hypothetical protein FLJ20392
	112888 T03872	AW195317 Hs.107716	hypothetical protein FLJ22344
	131863 T10072	Al656378 Hs.33461	ESTs
	112911 T10080	AW732747 Hs.13493	like mouse brain protein E46
25	132215 T10132	AL035703 Hs.4236	KIAA0478 gene product
	112931 T15343	T02966 Hs.167428	ESTs
	112984 T23457	T16971 Hs.289014	ESTs, Weakly similar to A43932 mucin 2 p
	112998 T23555	H11257 Hs.22968	Homo sapiens done IMAGE:451939, mRNA se
	133376 T23670	BE618768 Hs.7232	acetyl-Coenzyme A carboxylase alpha
30	113026 T23948	AA376654	eukaryotic translation initiation factor
	113070 T33464	AB032977 Hs.6298	KIAA1151 protein
	410781 T34413	Al375672 Hs.165028	ESTs
	113074 T34611	AK001335 Hs.31137	protein tyrosine phosphatase, receptor t
25	113095 T40920	AA828380 Hs.126733	ESTS
35	113179 T55182	BE622021 Hs.152571	ESTs, Highly similar to IGF-II mRNA-bind
	113337 T77453	T77453 Hs.302234	ESTs
	113421 T84039	Al769400 Hs.189729	ESTs
	113454 T86458 113481 T87693	Al022166 Hs.16188	ESTs
40	453345 T89350	T87693 Hs.204327 AA302862 Hs.90063	EST neurocalcin delta
40	113557 T90945	H66470 Hs.16004	ESTs
	113559 T90987	T79763 Hs.14514	ESTS
	113589 T91863	Al078554 Hs.15682	ESTs
	113591 T91881	T91881 Hs.200597	KIAA0563 gene product
45	113619 T93783	R08665 Hs.17244	hypothetical protein FLJ13605
	113683 T96687	AB035335 Hs.144519	T-ceil leukemia/lymphoma 6
	113692 T96944	AL360143 Hs.17936	DKFZP434H132 protein
	113702 T97307	T97307	gb:ye53h05.s1 Soares fetal liver spleen
	113717 T97764	T99513 Hs.187447	ESTs
50	113824 W48817	Al631964 Hs.34447	ESTs
	113840 W58343	R72137 Hs.7949	DKFZP586B2420 protein
	113844 W59949	Al369275 Hs.243010	Homo sapiens cDNA FLJ14445 fis, clone HE
•	113902 W74644	AA340111 Hs.100009	acyl-Coenzyme A oxidase 1, palmitoyl
	113904 W74761	AF125044 Hs.19196	ubiquitin-conjugating enzyme HBUCE1
55	113905 W74802	R81733 Hs.33106	ESTs
	113931 W81205	BE255499 Hs.3496	hypothetical protein MGC15749
	113932 W81237	AA256444 Hs.126485	hypothetical protein FLJ12604; KIAA1692
	131965 W90146	W79283 Hs.35962	ESTs
60	114035 W92798	W92798 Hs.269181	ESTs
60	114106 Z38412	AW602528	gb:RC5-BT0562-260100-011-A02 BT0562 Homo
	457308 Z38709	Al416988 Hs.238272	inositol 1,4,5-triphosphate receptor, ty
	114161 Z38904	BE548222 Hs.299883	hypothetical protein FLJ23399
	424949 Z39103	AF052212 Hs.153934	core-binding factor, runt domain, alpha
65	457548 Z39930 128937 Z39939	AW069534 Hs.279583	CGI-81 protein ESTs, Weakly similar to ALU1_HUMAN ALU S
05	432554 Z40012	AA251380 Hs.10726	NCK-associated protein 1
	114277 Z40377	Al479813 Hs.278411	
	114304 Z40820	AI052229 Hs.25373 AI934204 Hs.16129	ESTs, Weakly similar to T20410 hypotheti ESTs
	114364 Z41680	AL117427 Hs.172778	Homo sapiens mRNA; cDNA DKFZp566P013 (fr
70	432620 AA005112	AA777749 Hs.5978	LIM domain only 7
	129034 AA005432	AA481157 Hs.108110	DKFZP547E2110 protein
	131881 AA010163	AW361018 Hs.3383	upstream regulatory element binding prot
	332421 AA026356	Al909968 Hs.108106	transcription factor
	114465 AA026901	BE621056 Hs.131731	hypothetical protein FLJ11099
75	451271 AA036867	AK001644 Hs.26156	hypothetical protein FLJ10782
	332498 AA044644	AA303661	lymphocyte-specific protein 1

	104555 11040100	11045470 11-000004	Ode40 effectes entitle 2
	431555 AA046426	Al815470 Hs.260024	Cdo42 effector protein 3 Homo sapiens cDNA: FLJ23020 fis, clone L
	132944 AA054515 114618 AA084162	T96641 Hs.6127 AW979261 Hs.291993	ESTs
	332509 AA085749	AA128376 Hs.153884	ATP binding protein associated with cell
5	114648 AA101056	AA101056	gb:zn25b03.s1 Stratagene neuroepithelium
,	114658 AA102746	AA102383 Hs.249190	tumor necrosis factor receptor superfami
	132456 AA114250	AB011084 Hs.48924	KIAA0512 gene product; ALEX2
	450847 AA126561	NM_003155Hs.25590	stanniocalcin 1
	132225 AA128980	AA128980	gb:zo09a11.s1 Stratagene neuroepithelium
10	437197 AA129757	W38586	guanine nucleofide binding protein (G pr
	114709 AA129921	AA397651 Hs.301959	profine synthetase co-transcribed (bacte
	456926 AA133331	AB018284 Hs.158688	KIAA0741 gene product
	114750 AA135958	AA887211 Hs.129467	ESTs
	426806 AA136524	T19228 Hs.172572	hypothetical protein FLJ20093
15	114763 AA147044	AA810755 Hs.102500	hypothetical protein dJ511E16.2
	114767 AA148885	AI859865 Hs.154443	minichromosome maintenance deficient (S.
	114774 AA150043	AV656017 Hs.184325	CGI-76 protein
	129388 AA151621	AA662477 Hs.110964	hypothetical protein FLJ23471
20	457742 AA155743	BE561824 Hs.273369	uncharacterized hematopoletic stem/proge
20	456200 AA156335	AA768242 Hs.80618	hypothetical protein
	130207 AA156336	AF044209 Hs.144904	nuclear receptor co-repressor 1
	114798 AA159181	AA159181 Hs.54900 Z19448 Hs.131887	serologically defined colon cancer antig ESTs, Weakly similar to T24396 hypotheti
	114800 AA159825 114828 AA234185	AA252937 Hs.283522	Homo sapiens mRNA; cDNA DKFZp434J1912 (f
25	114846 AA234929	BE018682 Hs.166196	ATPase, Class I, type 8B, member 1
2,5	114848 AA234935	BE614347 Hs.169615	hypothetical protein FLJ20989
	114902 AA236359	AW275480 Hs.39504	hypothetical protein MGC4308
	132271 AA236466	AB030034 Hs.115175	sterile-alpha motif and leucine zipper c
	114907 AA236535	N29390 Hs.13804	hypothetical protein dJ462O23.2
30	420170 AA236935	U43374 Hs.95631	Human normal keratinocyte mRNA
	132204 AA236942	AA235827 Hs.42265	ESTs
	114928 AA237018	AA237018 Hs.94869	ESTs
	132481 AA237025	W93378 Hs.49614	ESTs
25	114932 AA242751	AA971436 Hs.16218	KIAA0903 protein
35	314162 AA242760	BE041820 Hs.38516	Homo sapiens, done MGC:15887, mRNA, com
	131006 AA242763	AF064104 Hs.22116	CDC14 (cell division cycle 14, S. cerevi
	114935 AA242809	H23329 Hs.290880	ESTs, Weakly stmilar to ALU1_HUMAN ALU S
	408908 AA243133 .	BE296227 Hs.250822	serine/threonine kinase 15
40	437754 AA243495 114957 AA243706	R60366 Hs.5822 AW170425 Hs.87680	Homo sapiens cDNA: FLJ22120 fis, clone H ESTs
40	114974 AA250848	AW966931 Hs.302649	nucleosome assembly protein 1-like 1
	114977 AA250868	AW296978 Hs.87787	ESTs
	114995 AA251152	AA769266 Hs.193657	ESTs
	115005 AA251544	AI760825 Hs.153042	ESTs
45	417177 AA251792	NM_004458Hs.81452	fatty-acid-Coenzyme A ligase, long-chain
	115026 AA252144	AA251972 Hs.188718	ESTs
	115045 AA252524	AW014549 Hs.58373	ESTs
	115068 AA253461	AW512260 Hs.87767	ESTs
	133138 AA255522	AV657594 Hs.181161	Homo sapiens cONA FLJ14643 fis, clone NT
50	332668 AA255522	AV657594 Hs.181161	ESTs
	115114 AA256468	AA527548 Hs.7527	small fragment nuclease
	129584 AA256528	AV656017 Hs.184325	CGI-76 protein
	115137 AA257976	AW968304 Hs.56156	ESTs
55	417187 AA258296	AB011151 Hs.334659	hypothetical protein MGC14139
33	115166 AA258409 115167 AA258421	AF095727 Hs.287832 AA749209 Hs.43728	myelin protein zero-like 1 hypothetical protein
	436719 AA262077	Y11192 Hs.5299	aldehyde dehydrogenase 5 family, member
	115239 AA278650	BE251328 Hs.73291	hypothetical protein FLJ10881
	115243 AA278766	AA806600 Hs.116665	KIAA1842 protein
60	428419 AA280791	U49436	KIAA1856 protein
• •	115322 AA280819	L08895 Hs.78995	MADS box transcription enhancer factor 2
	413303 AA280828	AW836130 Hs.75277	hypothetical protein FLJ13910
	115372 AA282195	AW014385 Hs.88678	ESTs, Weakly similar to Unknown [H.saple
	409962 AA283127	U82671 Hs.57698	Target CAT
65	130269 AA284694	F05422 Hs.168352	nucleoporin-like protein 1
	456570 AA291137	AA286914 Hs.183299	ESTs
	332675 AA291708	BE439944	ESTs
	407864 AA293495	AF069291 Hs.40539	chromosome 8 open reading frame 1
70	115536 AA347193	AK001468 Hs.62180	anillin (Drosophila Scraps homolog), act
70	408799 AA398474	AA059412 Hs.47986	hypothetical protein MGC10940
	115575 AA398512	AA393254 Hs.43619	ESTS .
	115601 AA400277	AA148984 Hs.48849	ESTs, Weakly similar to ALU4_HUMAN ALU S
	434428 AA400896	D14540 Hs.199160	myeloid/lymphoid or mixed-lineage leukem
75	115683 AA410345 115715 AA416733	AF255910 Hs.54650 BE395161 Hs.1390	junctional adhesion molecule 2 proteasome (prosome, macropain) subunit,
15	132952 AA425154	Al658580 Hs.61426	Homo saplens mesenchymal stem cell prote
	132332 744123134	130000 1301420	House achiera incacimilanes areas ocu biore

		AA426573		Hs.41135	endomucin-2
	409124	AA431418	AW292809	Hs.50727	N-acetylglucosaminidase, alpha- (Sanfill
	115895	AA436182	AB033035	Hs.51965	KIAA1209 protein
	458073	AA437099	AA192669	Hs.45032	ESTs
5		AA446585	AI636361		hypothetical protein MGC10702
•		AA446887	AJ745379		ESTs
		AA447224		Hs.238944	hypothetical protein FLJ10631
	_	AA447709		Hs.268115	ESTs, Wealty similar to T08599 probable
4.0		AA453624	AA252468	Hs.1098	DKFZp434J1813 protein
10	446730	AA455044	BE384932	Hs.64313	ESTs, Wealty similar to AF257182 1 G-pro
	116095	AA456045	AA043429	Hs.62618	ESTs
	426856	AA460454	R19768	Hs.172788	ALEX3 protein
		AA476494		Hs.172788	ALEX3 protein
		AA476738		Hs.326740	
15					hypothetical protein MGC10947
15		AA481422	D14041	Hs.347340	H-2K binding factor-2
		AA482595		Hs.55189	hypothetical protein
		AA485084	AW157022	Hs.343551	hypothetical protein FLJ22584
	116274	AA485431	Al129767	Hs.182874	guanine nucleotide binding protein (G pr
	426002	AA489638	BE514376	Hs.165998	PAI-1 mRNA-binding protein
20	116331	AA491000	N41300	Hs.71616	Homo sapiens mRNA; cDNA DKFZp586N1720 (f
		AA491250		Hs.203963	hypothetical protein FLJ10339
		AA505133		Hs.279905	done HQ0310 PRO0310p1
		AA598447		Hs.85951	exportin, tRNA (nuclear export receptor
25		AA599243	T86558	Hs.75113	general transcription factor IIIA
25		AA599574		3Hs.65370	lipase, endothelial
	134531	AA600153	Al742845	Hs.110713	DEK oncogene (DNA binding)
	116417	AA609309	AW499664	ļ.	Human clone 23826 mRNA sequence
	116429	AA609710	AF191018	Hs.279923	putative nucleofide binding protein, est
	116439	AA610068	AA251594	Hs.43913	PIBF1 gene product
30		AA621399	R80137	Hs.302738	Homo sapiens cDNA: FLJ21425 fis, clone C
••		AA621752		Hs.178761	26S proteasome-associated pad1 homolog
	409633			Hs.55200	ESTs
		D12160	D12160		
		-		Hs.249212	polymerase (RNA) III (DNA directed) (155
25		D19708		Hs.169531	ESTs
35	414964			Hs.333402	hypothetical protein MGC12760
	116571		D45652	Hs.211604	gb:HUMGS02848 Human adult lung 3' direct
	451522	D60208	BE565817	Hs.26498	hypothetical protein FLJ21657
	421919	D80504	AJ224901	Hs.109526	zinc finger protein 198
	116643	F03010	Al367044	Hs.153638	myeloid/lymphoid or mixed-lineage leukem
40	116661	F04247	R61504		gb:yh16a03.s1 Soares infant brain 1NIB H
	116715			Hs.170263	tumor protein p53-binding protein, 1
	116729			Hs.115823	ribonuclease P, 40kD subunit
	318709		R52576	Hs.285280	Homo sapiens cDNA: FLJ22096 fis, clone H
15	418999		NM_00012		erythropoletin receptor
45	116773		AI823410	Hs.343581	karyopherin alpha 1 (importin alpha 5)
	116780		H22566	Hs.63931	ESTs
	453884		AA355925	Hs.36232	KIAA0186 gene product
	116819	H53073	H53073	Hs.93698	EST
	427278	H56559	AL031428	Hs.174174	KIAA0601 protein
50	407833	H57957	AW955632	Hs.66666	ESTs, Weakly similar to S19560 proline-r
	116844		H64938	Hs.337434	ESTs, Weakly similar to A46010 X-linked
	116845			Hs.348148	gb:ns44f05.s1 NCI_CGAP_Alv1 Homo saplens
	116892		AI573283		ESTs
	116925			Hs.260603	ESTs, Moderately similar to A47582 B-cel
55			H73110		
"	116981		N29218	Hs.40290	ESTs
	453133		AC005757		hypothetical protein
	117031		H88353	Hs.347265	gb:yw21a02.s1 Morton Fetal Cochlea Homo
	117034		U72209		YY1-associated factor 2
	431129	H88675	AL137751	Hs.263671	Homo sapiens mRNA; cDNA DKFZp434l0812 (f
60	417861	H93708	AA334551		sperm specific antigen 2
	117280	N22107	M18217	Hs.172129	Homo saplens cDNA: FLJ21409 fis, clone C
	117344		R19085	Hs.210705	Homo sapiens cDNA FLJ13182 fis, clone NT
	117422		Al355562	Hs.43880	ESTs, Weakly similar to A46010 X-linked
	117475		N30205	Hs.93740	ESTs, Weakly similar to 138022 hypotheti
65	117475			Hs.44203	ESTs, Weakly Similar to 138022 hypotheti
UJ			N30621		
	117937		AF044209		nuclear receptor co-repressor 1
	130207		AF044209	Hs.144904	nuclear receptor co-repressor 1
	117549		N33390	Hs.44483	EST
	117683	N40180	N40180		gb:yy44d02.s1 Soares_multiple_sclerosis_
70	117710	N45198	N45198	Hs.47248	ESTs, Highly similar to similar to Cdc14
	117791	N48325	N48325	Hs.93956	EST
	117822		AA706282		ESTs
	422544		AB018259	Hs.118140	KIAA0716 gene product
	117895		AW450348	Hs 93998	ESTs, Highly similar to SORL_HUMAN SORTI
75	452259		AA317439	He 28707	
15			MAJ 17439	Un CADO4	signal sequence receptor, gamma (translo
	133057	NJ3143	AA465131	IDS.04001	Homo sapiens done 25218 mRNA sequence

	119102	N55326	A A 404722	Hs.184134	ESTs .
		N55493	N55493	FIS. 104 134	
					gb:yv50c02.s1 Soares fetal liver spleen
		N57493	N57493	11-040400	gb:yy54c08.s1 Soares_multiple_sclerosis_
5		N62955	N62955	Hs.316433	Homo saplens cDNA FLJ11375 fis, clone HE
3		N63520	N63520		gb:yy62f01.s1 Soares_multiple_sclerosis_
		N63604	BE327311		HT021
		N64166		Hs.173859	frizzled (Drosophila) homolog 7
		N64168	Al183838	Hs.48938	hypothetical protein FLJ21802
••		N64191	N46114	Hs.29169	hypothetical protein FLJ22623
10	118475	N66845	N66845		gb:za46c11.s1 Soares fetal liver spleen
	118491	N67135	AV647908	Hs.90424	Homo sapiens cDNA: FLJ23285 fis, clone H
	118500	N67295	W32889	Hs.154329	ESTs
	118584	N68963	AW136928		gb:UI-H-BI1-adp-d-08-0-UI.s1 NCI_CGAP_Su
	456647	N69331	Al252640	Hs.110364	peptidytorolyl isomerase C (cyclophilin
15	118661	N70777	AL137554	Hs.49927	protein kinase NYD-SP15
	118684	N71364	N71313	Hs.163986	Homo sapiens cDNA: FLJ22765 fis, clone K
	118689	N71545	AW390601	Hs.184544	Homo sapiens, done IMAGE:3355383, mRNA,
	118690	N71571	N71571	Hs.269142	ESTs
	118766	N74456	N74456	Hs.50499	EST
20	118793	N75594	N75594	Hs.285921	ESTs, Moderately similar to T47135 hypot
		N79035	AI668658	Hs.50797	ESTs
		N80279	AL035364		hypothetical protein
		N91797	AW452696		myosin phosphatase, target subunit 2
		N92454	AW580922		karyopherin (importin) beta 1
25		N94581	AW191962		collagen, type VIII, alpha 2
		N94746	N94746	Hs.274248	hypothetical protein FLJ20758
		N98238	N98238	Hs.55185	ESTs :
		R02384	Al160570	Hs.252097	pregnancy specific beta-1-glycoprotein 6
		R16833	R16833	Hs.53106	ESTs, Moderately similar to ALU1_HUMAN A
30		R41828	R10674	110.00100	CSR1 protein
-		R43203	T02865	Hs.328321	EST
		R46395 ·			hypothetical protein
		R58863	R58863	Hs.91815	ESTs
		R78248	AW995911		hypothetical protein FLJ23399
35		T11483	T11483		gb:CHR90049 Chromosome 9 exon Homo saple
		T16896	AI692322	Hs.65373	ESTs, Weakly similar to T02345 hypotheti
		T23820	NM_001241		cyclin T2
		T30222	T10077	Hs.13453	hypothetical protein FLJ14753
		W15275		Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
40		W38194	W38194		Empirically selected from AFFX single pr
		W42414	AW081883	Hs.211578	Homo sapiens cDNA: FLJ23037 fis, clone L
	419445	W49632	AA884471		Human clone 23908 mRNA sequence
	119650	W57613	R82342	Hs.79856	ESTs, Weakly similar to S65657 alpha-1C-
	119654	W57759	W57759		gb:zd20g11.s1 Soares_fetal_heart_NbHH19W
45	119683	W61118	W65379	Hs.57835	ESTs
	119694	W65344		Hs.57847	ESTs, Moderately similar to ICE4_HUMAN C
	119718	W69216	W69216	Hs.92848	ESTs
	410365	W69379	AI287518		Homo sapiens mRNA; cDNA DKFZp586D0923 (f
	119938	W86728	AW014862	Hs.58885	ESTs
50	120128	Z38499	BE379320	Hs.91448	MKP-1 like protein tyrosine phosphatase
	120130	Z38630	AA045767	Hs.5300	bladder cancer associated protein
	120148	Z39494	F02806	Hs.65765	ESTs
	120155	Z39623	Z39623	Hs.65783	ESTs
	451979		F06972	Hs.27372	BMX non-receptor tyrosine kinase
55	120183	Z40174	AW082866	Hs.65882	ESTs
	120184	Z40182		Hs.65885	EST
	120211	Z40904	Z40904	Hs.66012	EST
		AA166965	AW959615		ESTs .
	120247	AA167500	AA167500		EST
60	120254	AA169599	W90403	Hs.111054	ESTs
	120259	AA171724	AW014786	Hs.192742	hypothetical protein FLJ12785
	120260	AA171739	AK000061	Hs.101590	hypothetical protein
	120275	AA177105	AA177105	Hs.78457	solute carrier family 25 (mitochondrial
	120284	AA182626	AA179656		gb:zp54e11.s1 Stratagene NT2 neuronal pr
65	417735	AA186324	AA188175	Hs.82506	KIAA1254 protein
		AA192099	AJ236885		zinc finger protein 148 (pHZ-52)
	120302	AA192173	AA837098	Hs.269933	ESTs
		AA192415	Al216292	Hs.96184	ESTs
	120305	AA192553	AW295096	Hs.101337	uncoupling protein 3 (mitochondrial, pro
70	120319	AA194851	T57776	Hs.191094	ESTs
	408729	AA195520	AA195764	Hs.72639	ESTs
		AA196300	AA196300		hypothetical protein RG083M05.2
		AA196549		Hs.6592	Homo sapiens, done IMAGE:2961368, mRNA,
		AA196721	AK000292		hypothetical protein FLJ20285
75		AA196979	AA923278	Hs.290905	ESTs, Wealdy similar to protease [H.sapi
	120340	AA206828	AA206828		gb:zq80b08.s1 Stratagene hNT neuron (937

	417122 AA207123	Al906291 Hs.81234	immunoglobulin superfamily, member 3
	131522 AA214539	Al380040 Hs.239489	TIA1 cytotoxic granule-associated RNA-bi
	421787 AA226914	AA227068 Hs.108301	nuclear receptor subfamily 2, group C, m
	120375 AA227260	AF028706 Hs.111227	Zic family member 3 (odd-paired Drosophi
5	120376 AA227469	AA227469	gb:zr18a07.s1 Stratagene NT2 neuronal pr
	120390 AA233122	AA837093 Hs.111460	calcium/calmodulin-dependent protein kin
	410804 AA233334	U64820 Hs.66521	Machado-Joseph disease (spinocerebellar
	434223 AA233347	Al825842 Hs.3776	zinc finger protein 216
	312771 AA233714	AA018515 Hs.264482	Homo saplens mRNA; cDNA DKFZp761A0411 (i
10	120396 AA233796	AA134006 Hs.79306	eukaryotic translation initiation factor
10	120409 AA235050	AA235050	gb:zs38e04.s1 Soares_NhHMPu_S1 Homo sapi
	120414 AA235704	AW137156 Hs.181202	
	120420 AA236031	Al128114 Hs.112885	hypothetical protein FLJ10038 spinal cord-derived growth factor-B
	120422 AA236352		•
· 15	419326 AA236390		hypothetical protein DKFZp434N1928 ESTs
13	120423 AA236453		
	120435 AA243370	AA236453 Hs.18978 AA243370 Hs.96450	Horno sapiens cDNA: FLJ22822 fis, clone K
			EST
	120453 AA250947	AA250947 Hs.170263	tumor protein p53-binding protein, 1
20	120455 AA251083	AA251720 Hs.104347	ESTs, Wealdy similar to ALUC_HUMAN !!!!
20	120456 AA251113	AA488750 Hs.88414	BTB and CNC homology 1, basic leucine zi
	120473 AA251973	AA251973 Hs.269988	ESTs
	128922 AA252023	Al244901 Hs.9589	ubiquilin 1
	120477 AA252414	AA252414 Hs.43141	DKFZP727C091 protein
25	120479 AA252650	AF006689 Hs.110299	mitogen-activated protein kinase kinase
25	120488 AA255523	AW952916 Hs.63510	KIAA0141 gene product
	120510 AA258128	Al796395 Hs.111377	ESTs
	120527 AA262105	AA262105 Hs.4094	Homo saplens cDNA FLJ14208 fis, clone NT
	120528 AA262107	Al923511 Hs.104413	ESTs ,
	120529 AA262235	Al434823 Hs.104415	ESTs
30	120541 AA278298	W07318 Hs.240	M-phase phosphoprotein 1
	120544 AA278721	BE548277 Hs.103104	ESTs
	120562 AA280036	BE244580 Hs.342307	hypothetical protein FLJ10330
	120569 AA280648	AA807544 Hs.24970	ESTs, Weakly similar to B34323 GTP-bindi
	120571 AA280738	AB037744 Hs.34892	KIAA1323 protein
35	120572 AA280794	H39599 Hs.294008	ESTs
	129434 AA280837	AW967495 Hs.186644	ESTs
	130529 AA280886	AA178953 Hs.309648	gb:zp39e03.s1 Stratagene muscle 937209 H
	120575 AA280934	AW978022 Hs.238911	hypothetical protein DKFZp762E1511; KIAA
	409339 AA281535	AB020686 Hs.54037	ectonucleotide pyrophosphatase/phosphodi
40	120591 AA281797	AF078847 Hs.191356	general transcription factor IIH, polype
	120593 AA282047	AA748355 Hs.193522	ESTs
	430275 AA283002	Z11773 Hs.237786	zinc finger protein 187
	440303 AA283709	AA306166 Hs.7145	calpain 7
	120609 AA283902	AW978721 Hs.266076	ESTs, Weakly similar to A46010 X-linked
45	409702 AA284108	Al752244	eukaryotic translation elongation factor
	456870 AA284109	Al241084 Hs.154353	nonselective sodium potassium/proton exc
	132614 AA284371	AA284371 Hs.118064	similar to rat nuclear ubiquitous casein
	458750 AA284744	AA115496 Hs.336898	Homo saplens, Similar to RIKEN cDNA 1810
	135376 AA284784	BE617856 Hs.99756	mitochondrial ribosome recycling factor
50	120621 AA284840	AW961294 Hs.143818	hypothetical protein FLJ23459
•	452279 AA286844	AA286844 Hs.61260	hypothetical protein FLJ13164
	332484 AA287032	AW172431 Hs.13012	ESTs
	120644 AA287038	Al869129 Hs.96616	ESTs
	120660 AA287546	AA286785 Hs.99677	ESTs
55	135370 AA287553	BE622187 Hs.99670	ESTs, Weakly similar to 138022 hypotheti
-	120661 AA287556	AA287556 Hs.263412	ESTs, Weakly similar to ALUB_HUMAN !!!!
	429828 AA287564	AB019494 Hs.225767	
		AF015592 Hs.28853	IDN3 protein
•	452291 AA291015		CDC7 (cell division cycle 7, S. cerevisi
60	120699 AA291716	A1683243 Hs.97258	ESTs, Moderately similar to S29539 ribos
UU	100690 AA291749	AA383256 Hs.1657	estrogen receptor 1
	120726 AA293656	AA293655 Hs.21198	ESTs
	120737 AA302430	AL049176 Hs.82223	chordin-like
	120745 AA302809	AA302809	gb:EST10426 Adipose tissue, white I Homo
45	443574 AA302820	U83993 Hs.321709	purinergic receptor P2X, ligand-gated lo
65	120750 AA310499	Al191410 Hs.96693	ESTs, Moderately similar to 2109260A B c
	120761 AA321890	AA321890	branched chain keto acid dehydrogenase E
	120768 AA340589	AA340589 Hs.104560	EST
	120769 AA340622	Al769467 Hs.9475	ESTs
	135232 AA342457	AL038812 Hs.96800	ESTs, Moderately similar to ALU7_HUMAN A
70	120793 AA342864	AA342864 Hs.96812	ESTs
	120796 AA342973	Al247356 Hs.96820	ESTs
	120809 AA346495	AA346495	gb:EST52657 Fetal heart II Homo saptens
	332633 AA347573	AL120071 Hs.48998	fibronedin leudne rich transmembrane p
	120825 AA347614	Al280215 Hs.96885	ESTs
75	120827 AA347717	AA382525 Hs.132967	Human EST clone 122887 mariner transposo
	120839 AA348913	AA348913	gb:EST55442 Infant adrenal gland II Homo
			g

	120850 AA349647	AA349647	Hs.96927	Homo saplens cDNA FLJ12573 fis, clone NT
	120852 AA349773		Hs.191564	ESTs
	128852 AA350541	R40622	Hs.106601	ESTs
. 5	135240 AA357159 120870 AA357172		Hs.96986	EST
	120894 AA370132		Hs.292581 Hs.97063	ESTs, Moderately similar to ALU1_HUMAN A ESTs
	435737 AA370472		Hs.173202	I-kappa-B-interacting Ras-like protein 1
	120897 AA370867		Hs.97079	ESTs, Moderately similar to AF174605 1 F
	120915 AA377296	AL135556	Hs.97104	ESTs
10	120935 AA383902		Hs.97177	ESTs, Weakly similar to ALU1_HUMAN ALU S
	120936 AA385934		Hs.97184 Hs.97186	EST, Highly similar to (defline not avai
	120937 AA386255 120938 AA386260		Hs.104632	EST .
	417632 AA386266	R20855	Hs.5422	glycoprotein M6B
15	120960 AA398014		Hs.104684	EST
	120985 AA398222	Al219896	Hs.97592	ESTs
	120988 AA398235	AA398235		ESTs
	121008 AA398348 121029 AA398482	AA398348 AA398482	Hs.130546	Human DNA sequence from clone RP11-251J8 EST
20	121032 AA398504		Hs.161798	ESTs
-0	121033 AA398505		Hs.97360	ESTs
	121034 AA398507		Hs.271623	nucleoporin 50kD
	121035 AA398523		Hs.210579	ESTs
25	121058 AA398625	AA398625		ESTs
23	121060 AA398632 121061 AA398633	AA398632 AA393288		ESTs ESTs
	121091 AA398894	AA398894		ESTs, Moderately similar to ALU8_HUMAN A
	121092 AA398895	AA398895		EST
••	121094 AA398900	AA402505		gb:zt62h10.r1 Soares_testis_NHT Homo sap
30	121096 AA398904		Hs.332690	ESTs
	121115 AA399122 121121 AA399371		Hs.104682 Hs.189095	ESTs, Weakly similar to mitochondrial ci similar to SALL1 (sal (Drosophila)-like
	121121 AA399373	Al126713		ESTs, Highly similar to T00337 hypotheti
	121125 AA399441	AL042981		KIAA1201 protein
35	121151 AA399636	AA399636	Hs.143629	ESTs
	121153 AA399640	AA399640		ESTs
	121163 AA399680	Al676062	Hs.111902	ESTs
	121176 AA400080 121192 AA400262	AL121523 AA400262	Hs.190093	ESTs ESTs
40	121223 AA400725	AJ002110	Hs.97169	ESTs, Weakly similar to dJ667H12.2.1 [H.
	121227 AA400748	AA400748	Hs.97823	Homo sapiens mRNA; cDNA DKFZp434D024 (fr
	121231 AA400780	AA814948		ESTs, Weakly similar to ALUC_HUMAN !!!!
	121278 AA401631 121279 AA401688	AA037121	Hs.98518 Hs.177996	Homo sapiens cDNA FLJ11490 fis, clone HE ESTs
45	121282 AA401695	AA401695		ESTs
	121299 AA402227	AA402227		tropomodulin 3 (ubiquitous)
	121301 AA402329	NM_00620	2Hs.89901	phosphodiesterase 4A, cAMP-specific (dun
	121302 AA402398		Hs.325520	LAT1-3TM protein
50	121304 AA402449 121305 AA402468	AA293863	Hs.9/316 Hs.291557	EST ESTs
50	134721 AA403268	AK000112		hypothetical protein FLJ20105
	121323 AA403314	AA291411		ESTs
	121324 AA404229	AA404229	Hs.97842	EST
EE	444422 AA404260	Al768623	Hs.108264	ESTs
55	131074 AA404271 121344 AA405026	U16125	Hs.181581 Hs.193754	glutamate receptor, ionotropic, kainate
	121348 AA405182		Hs.97973	ESTs ESTs
	121350 AA405237	AA405237		gb:zt06e10.s1 NCI_CGAP_GCB1 Homo sapiens
	121400 AA406061	AA406061		EST
60	121402 AA406063	AA406063		ESTs
	121403 AA406070 121408 AA406137	AA406070 AA406137		EST EST
	121431 AA406335		Hs.176731	ESTs
_	121471 AA411804		Hs.261575	ESTs
65	121474 AA411833		Hs.188760	ESTs, Highly similar to Trad [H.sapiens]
	121526 AA412219	AW665325		ESTs
	121530 AA412259 121558 AA412497	AA778658 AA412497	HS.98122	ESTs gb:zt95g12.s1 Soares_testis_NHT Homo sap
	121559 AA412498	Al192044	Hs.104778	ESTs
70	121584 AA416586	AI024471		ESTs
	121609 AA416867	AA416867		EST
	121612 AA416874	AA416874		ESTs
	121737 AA421133 121740 AA421138		Hs.104671 Hs.143835	erythrocyte transmembrane protein EST
75	436032 AA422079		Hs.109276	latexin protein
	121784 AA423837	T90789	Hs.94308	RAB35, member RAS oncogene family

	121902 A	A 424229	Al251870	U. 400000	ECT
	121802 A 121803 A		Al338371	Hs.188898 Hs.157173	ESTs ESTs
	135286 A		AW023482		ESTs
	332778 A		AW023482		ESTs
5	121806 A		AA424313		ESTs
_	129517 A			Hs.112237	ESTs
	121845 A	A425734	AI732692	Hs.165066	ESTs, Moderately similar to ALU2_HUMAN A
	121853 A		AA425887		hypothetical protein FLJ14303
10	121891 A		AA426456	Hs.98469	ESTs
10	121895 A		AA427396	D- 50404	gb:zw33a02.s1 Soares ovary tumor NbHOT H
	121899 A 121917 A		R55341 AA406397	Hs.50421 Hs.139425	KIAA0203 gene product ESTs
	121918 A			Hs.184175	chromosome 2 open reading frame 3
	121919 A		AA428281		EST
15	121941 A	A428865	AA428865		ESTs
	121942 A		AW452701	Hs.293237	ESTs
	121970 A		AA429666		EST
	121993 A		AW297880		ESTs
20	418706 A 122022 A		U73524 AA431293	Hs.87465	ATP/GTP-binding protein
20	122050 A		Al453076	115.307 10	ESTs, Moderately similar to T42650 hypot ELAV (embryonic lethal, abnormal vision,
	122051 A		AA431492	Hs.98742	EST
	122055 A		AA431732		EST
	122105 A		AW241685	Hs.98699	ESTs
25	122125 A		AK000492		hypothetical protein
	135235 A			Hs.266195	ESTs
	122162 A/ 422072 A/		AA628233	Hs.111138	cytochrome P450, subfamily XIX (aromatiz KIAA0712 gene product
	415106 A		U40763	Hs.77965	peptidyl-prolyl isomerase G (cyclophilin
30	122186 A			Hs.104673	ESTs
	122235 A	A436475	AA436475	Hs.112227	membrane-associated nucleic acid binding
	412970 A			Hs.177534	dual specificity phosphatase 10
	419288 A/ 122310 A/		AA256106		ESTS
35	122310 AV		AW192803 BE465894		ESTs, Weakly similar to S65824 reverse t ESTs, Weakly similar to LB4D_HUMAN NADP-
-	122382 A		AA446440		ESTs
	122425 A		AB007859		KIAA0399 protein
	122431 A		AA447398		ESTs
40	122450 AA 426284 AA		AA447643		hypothetical protein DKFZp434F1819
70	122477 A		AJ404468 AA448226		dynein, axonemal, heavy polypeptide 9 ESTs
	122500 A		AA448825		ESTs
	122522 A		AA299607		ESTs
45	122536 A		AF060877		regulator of G-protein signalling 20
45	122538 A		AA450211		ESTS
	122540 AA 122560 AA		AA476741 AW392342		ESTs, Weakly similar to A43932 mucin 2 p centrosomal P4.1-associated protein; unc
	421919 A		AJ224901		zinc finger protein 198
	122562 A		AA452156		gb:zx29c03.s1 Soares_total_fetus_Nb2HF8_
50	122585 A		Al681654	Hs.170737	hypothetical protein FLJ23251
	122608 A		AA453525	Hs.143077	ESTs
	122635 AA 122636 AA		AA454085 AW651706	Un 00540	gb:zx33a08.s1 Soares_total_fetus_Nb2HF8_
	122653 A		AW009166		hypothetical protein FLJ14007 ESTs
55	122660 A			Hs.180069	nuclear respiratory factor 1
	122703 A		AA456323		ESTs
	122724 A		AA457395		ESTs
	122749 AA 122772 AA		AA458850		ESTs, Weakly similar to B34087 hypotheti
60	430242 A		AW117452 U66669	Hs.236642	ESTs 3-hydroxyisobutyryi-Coenzyme A hydrolase
00	429838 A		AW904907		hypothetical protein FLJ13409; KIAA1711
	122777 A	A4597 0 2	AK001022	Hs.214397	hypothetical protein FLJ10160 similar to
	135362 AA		AA978128		ESTs, Weakly similar to T17454 diaphanou
65	122798 AA		AW366286		splicing factor (CC1.3)
05	122837 AA 122860 AA		AA461509 AA464414		ESTs, Weakly similar to putative p150 [H gb:zx78g01.s1 Soares ovary tumor NbHOT H
	122861 A		AA335721	Hs.213628	ESTs
	122910 A	4470084	AA470084	Hs.98358	ESTs
	132899 AA		AA476606		SMAD in the antisense orientation
70	122967 AA	\ 478521	AA806187	Hs.289101	glucose regulated protein, 58kD
	422845 AA		AA317841		hypothetical protein MGC2752
	123009 AA 128917 AA		AA535244 AI365215		RAB2, member RAS oncogene family oncogene TC21
	123081 AA		Al815486		Homo sapiens cDNA FLJ20738 fis, clone HE
75	123133 A		AA487264		Homo sapiens mRNA; cDNA DKFZp667N064 (fr
	123184 AA		BE247767		KIAA0870 protein

	332467 AA489630	NM_014700Hs.119004	KIAA0665 gene product
	123233 AA490225	AW974175 Hs.151875	ESTs, Weakly similar to MAPB_HUMAN MICRO
	123234 AA490227	NM_001938Hs.16697	down-regulator of transcription 1, TBP-b
	123236 AA490255	AW968504 Hs.123073	CDC2-related protein kinase 7
5	123255 AA490890	AA830335 Hs.105273	ESTs
-	430015 AA490916	AW768399 Hs.106357	ESTs
	448892 AA490925	AF084535 Hs.22464	epilepsy, progressive myodonus type 2,
	123259 AA490955	Al744152 Hs.283374	ESTs. Wealty similar to CA15_HUMAN COLLA
	123284 AA495812	AA488988 Hs.293796	ESTs
10	123286 AA495824	AA495824 Hs.188822	ESTs, Weakly similar to A46010 X-linked
	123315 AA496369	AA496369	gb:zv37d10.s1 Soares ovary tumor NbHOT H
	457397 AA504125	AW969025 Hs.109154	ESTs
	433049 AA521473	AU076668 Hs.334884	SEC10 (S. cerevisiae)-like 1
	123421 AA598440	AA598440 Hs.291154	EST, Wealdy similar to I38022 hypothetic
15	123449 AA598899	AL049325 Hs.112493	Homo saplens mRNA; cDNA DKFZp564D036 (fr
	426981 AA599244	AL044675 Hs.173081	KIAA0530 protein
	409986 AA599694	NM_014777Hs.57730	KIAA0133 gene product
	123497 AA600037	AA765256 Hs.135191	ESTs, Weakly similar to unnamed protein
00	123604 AA609135	AA609135 Hs.293076	ESTs
20	123712 AA609684	AA609684	Homo sapiens cDNA: FLJ21543 fis, clone C
	123731 AA609839	AA609839 Hs.334437	gb:ae62f01.s1 Stratagene lung carcinoma
	123800 AA620423	AA620423 Hs.112862	EST
	123841 AA620747	AA620747 Hs.112896	ESTs
25	123929 AA621364	AA621364 Hs.112981	ESTs
25	123978 C20653	T89832 Hs.170278	ESTs
	133184 D20085	AA001021 Hs.6685	thyroid hormone receptor interactor 8
	132835 D20749	Z83844 Hs.5790	hypothetical protein dJ37E16.5
	435147 D51285	AL133731 Hs.4774	Homo sapiens mRNA; cDNA DKFZp761C1712 (f
30	128695 D59972	NM_003478Hs.101299	Cultin 5
30	124029 F04112	F04112 Hs.312553	gb:HSC2JH062 normalized infant brain cDN
	124057 F13604 449316 H01662	AA902384 Hs.73853 Al609045 Hs.321775	bone morphogenetic protein 2
	130973 H05135	Al638418 Hs.1440	hypothetical protein DKFZp434D1428 DEAD/H (Asp-Giu-Ala-Asp/His) box polypep
	124106 H12245	H12245	gb:ym17a12.r1 Soares infant brain 1NIB H
35	124136 H22842	H22842 Hs.101770	EST
55	124165 H30894	H30039 Hs.107674	ESTs
	429627 H43442	NM_015340Hs.2450	leucyl-tRNA synthetase, mitochondrial
	124178 H45996	BE463721 Hs.97101	putative G protein-coupled receptor
	129948 H69281	Al537162 Hs.263988	ESTs
40	452114 H69485	N22687 Hs.8236	ESTs
	124+0826254	H69899 H69899	ab:vu/uc12.s) Weizmann Uliactory Edithei
	124+D826254 129056 H70627	H69899 H69899 Al769958 Hs.108336	gb:yu70c12.s1 Weizmann Olfactory Epithel ESTs, Weakly similar to ALUE_HUMAN !!!!
			gb:yu/uc12.s1 Weizmann Oliactory Epithel ESTs, Weakly similar to ALUE_HUMAN !!!! Homo saplens clone FLB6914 PRO1821 mRNA,
	129056 H70627	AI769958 Hs.108336	ESTs, Weakly similar to ALUE_HUMAN !!!!
45	129056 H70627 427580 H73260	AI769958 Hs.108336 AK001507 Hs.44143	ESTs, Weakly similar to ALUE_HUMAN !!!! Homo saplens done FLB6914 PRO1821 mRNA,
45	129056 H70627 427580 H73260 426793 H77531	AI769958 Hs.108336 AK001507 Hs.44143 X89887 Hs.172350	ESTs, Wealdy similar to ALUE_HUMAN !!!! Homo saptens done FLB6914 PRO1821 mRNA, HIR (histone cell cycle regulation defec
45	129056 H70627 427580 H73260 426793 H77531 124274 H80552 129078 H80737 457658 H93412	AI769958 Hs.108336 AK001507 Hs.44143 X89887 Hs.172350 H80552 Hs.102249	ESTs, Wealdy similar to ALUE_HUMAN !!!! Homo saptens done FLB6914 PRO1821 mRNA, HIR (histone cell cycle regulation defec EST lysosomal presentins associated rhomboid-like pro
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50556065	129056 H70627 427580 H73260 426793 H77531 124274 H80552 129078 H80737 457658 H93412 124315 H94892 437712 H95643 124324 H95652 452933 H97146 132231 H99131 421877 H99462 443123 H99837 132963 N22140 420473 N22197 417381 N23756 130365 N24134 456610 N24195 439311 N26739 124383 N27098 124387 N27637 129341 N33090 419793 N35967 124433 N39069 124441 N46441 132338 N48270 436575 N48365 124466 N51316 408048 N51499 124483 N53976 124484 N54157 124485 N54300 124484 N54157 124485 N54300 1244484 N54831 129200 N59849	AI769958 Hs.108336 AK001507 Hs.44143 X89887 Hs.172350 H80552 Hs.102267 AW952124 Hs.102267 AW952124 Hs.13094 NM_005402Hs.288757 X04588 Hs.85844 H96552 Hs.159472 AW391423 Hs.288555 AA662910 Hs.42635 AW250380 Hs.109059 AA094538 Hs.272808 AA099593 Hs.34851 AL118782 Hs.300208 AF164142 Hs.82042 W56119 Hs.155103 AF172066 Hs.106346 BEZ70668 Hs.155103 AF172066 Hs.106346 BEZ70668 Hs.155103 AF172066 Hs.106346 BEZ70668 Hs.151945 NZ7098 Hs.102463 NZ7637 Hs.109019 AI193519 Hs.226396 AI364933 Hs.108463 AZ80319 Hs.286840 AW450481 Hs.161333 AA353868 Hs.182982 AI473114 R10084 Hs.113319 NM_007203Hs.42322 AI821780 Hs.179864 H66118 Hs.285520 AB040933 Hs.15420 NS4831 Hs.271381 N59849 Hs.13565	ESTs, Wealdy similar to ALUE_HUMAN IIII Homo sapiens clone FLB6914 PRO1821 mRNA, HIR (histone ceil cycle regulation defec EST lysosomal presenllins associated rhomboid-like pro v-ral simian leukemta viral oncogene hom neurotrophic tyrosine kinase, receptor, Homo sapiens cDNA: FLJ22224 fis, clone H Homo sapiens cDNA: FLJ22224 fis, clone H hypothetical protein DKFZp434K2435 mitochondrial ribosomal protein L12 putative transcription regulation nuclea epsllon-tubulin Sec23-interacting protein p125 solute carrier family 23 (nucleobase tra eukaryotic translation initiation factor retinoic acid repressible protein mitochondrial ribosomal protein L43 EST ESTs hypothetical protein FLJ11126 serine/threonine kinase 24 (Sta20, yeast PRO1575 protein ESTs golgin-67 ESTs kinesin heavy chain member 2 A kinase (PRKA) anchor protein 2 ESTs ESTs, Wealdy similar to 2109260A B cell KJAA1500 protein
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	124522	N62375	NECOTE	Un 402724	EOT
		N63138	N62375 AA903424	Hs.102731 Hs.6786	ESTs ·
		N63172	D54120	Hs.146409	cell division cycle 42 (GTP-binding prot
		N63787	BE296313		ESTs, Weakly similar to 138022 hypotheti
5		N68168	N68168		gb:za11c01.s1 Soares fetal liver spleen
		N68201	N68201		ESTs, Weakly similar to I38022 hypotheti
		N68300	N68300	Hs.138485	gb:za12g07.s1 Soares fetal liver spleen
		N68321 N69575	N68321 N69575	Hs.231500 Hs.102788	EST .
10		N75007		Hs.199009	protein containing CXXC domain 2
		N75542	A1680737		Homo sapiens cDNA FLJ11918 fis; clone HE
	128473	N90066	T78277	Hs.100293	O-linked N-acetytghucosamine (GlcNAc) tr
		N91246	AW582962	Hs.102897	CGI-47 protein
15	124652		W19407	Hs.3862	regulator of nonsense transcripts 2; DKF
15		N93214	AB002316		KIAA0318 protein
	124671 133054			Hs.102951 Hs.291079	Homo sapiens cDNA FLJ10495 fis, clone NT
	425266		J00077	Hs.155421	ESTs, Weakly similar to T27173 hypotheti alpha-fetoprotein
	124720		R05283	110,100121	gb:ye91c08.s1 Soares fetal liver spicen
20	124722		T97733	Hs.185685	ESTs
	128944		AL137586		anaphase-promoting complex subunit 7
	132965		AI248173	Hs.191460	hypothetical protein MGC12936
	426504			Hs.170160	RAB2, member RAS oncogene family-like
25	438828 124757		AL134275 H11368	Hs.141055	hypothetical protein DKFZp761F2014
20	124762		AA553722		Homo sapiens done 23758 mRNA sequence ESTs, Moderately similar to A46010 X-tin
	124773		R45154	Hs.338439	ESTs
	135266		R41179	Hs.97393	KIAA0328 protein
	427961			Hs.143134	ESTs
30	414303			7Hs.165263	early development regulator 2 (homolog o
	128540			Hs.328317	EST
	124785 124792		W38537 R44357	Hs.280740 Hs.48712	hypothetical protein MGC3040 hypothetical protein FLJ20735
	124793		R44519	115.407 12	gb:yg24h04.s1 Soares infant brain 1NIB H
35	124799		R45088		gb:yg38g04.s1 Soares infant brain 1NIB H
	124812	R47948	R47948	Hs.188732	ESTS
	124821		H87832	Hs.7388	kelch (Drosophila)-like 3
	424123		AW966158		Homo sapiens cDNA FLJ12789 fis, clone NT
40	124835 124845		R55241 R59585	Hs.101214 Hs.101255	EST ESTs
70	124847		W07701	Hs.304177	Homo sapiens clone FLB8503 PRO2286 mRNA,
	440630			Hs.239388	Human DNA sequence from done RP1-304B14
	124861		R67567	Hs.107110	ESTs
4.5	332503			5Hs.150956	exostoses (multiple)-like 1
45	124879		R73588	Hs.101533	ESTs
	124892 124906		Al970003 H75964	Hs.23756	hypothetical protein similar to swine ac
	124922		R93622	Hs.107815 Hs.12163	ESTs eukaryotic translation initiation factor
	124940		AF068846		heterogeneous nuclear ribonucleoprotein
50	124941		AI766661	Hs.27774	ESTs, Highly similar to AF161349 1 HSPC0
	124943	T02888	AW963279	Hs.123373	ESTs, Weakly similar to ALU1_HUMAN ALU S
	124947		T03170	Hs.100165	ESTs
	124954		AW964237		KIAA1548 protein
55	456862 410653		U55184 BE383768	Hs.154145	hypothetical protein FLJ11585 95 kDa retinoblastoma protein binding pr
55	418133		R43504	Hs.6181	ESTs
	440014		AW960782		ash2 (absent, small, or homeotic, Drosop
	131082		AI091121	Hs.246218	Homo sapiens cDNA: FLJ21781 fis, clone H
CO	124980		T40841	Hs.98681	ESTs
60	124984		BE313210	Hs.334798	eukaryotic translation elongation factor
	124991		T50116	N I - 000370	gbryb77c10.s1 Stratagene ovary (937217)
	457222 125000		NM_004477 T58615	Hs.203772 Hs.235887	FSHD region gene 1 ESTs
	132932		AW118826		Homo sapiens cONA: FLJ22783 fis, clone K
65 ·	444484		AK002126	Hs.11260	hypothetical protein FLJ11264
	125008		T91251		gb:yd60a10.s1 Soares fetal liver spleen
	125009		T64924	Hs.303046	ESTs
•	445384		T79136	Hs.127243	Homo sapiens mRNA for KIAA1724 protein,
70	125017		T68875	He 200404	gb:yc30f05.s1 Stratagene liver (937224)
70	125018 125020		T69027 T69981	Hs.269481	sex comb on midleg homolog 1
	437871		Al084813	Hs.114088	gb:yc19d03_r1 Stratagene lung (937210) H ESTs
	134204		AJ873257	Hs.7994	hypothetical protein FLJ20551
	125050		AW970209		ESTs
75	125052		T85104	Hs.222779	ESTs, Moderately similar to similar to N
	125054	180522	T80622	Hs.268601	ESTs, Wealtly similar to envelope [H.sapi

	405000 705000	T 0.000	
	125063 T85352	T85352	gb:yd82d01.s1 Soares fetal liver spleen
	125064 T85373	T85373 T86284	gb:yd82f07.s1 Soares fetal liver spleen
	125066 T86284 416507 T89579	,	gb:yd77b07.s1 Soares fetal liver spleen
5	125080 T90360	AL045364 Hs.79353	transcription factor Dp-1
,	125097 T94328	T90360 Hs.268620 AW576389 Hs.335774	ESTs, Highly straiter to ALU6_HUMAN ALU S
	125104 T95590	T95590	EST, Moderately similar to S65657 alpha-
	135107 T97257		gb:ye40a03.s1 Soares fetal liver spleen ESTs, Moderately similar to 138022 hypot
	423122 T97599	T97257 Hs.94560 AA845462 Hs.124024	
10	125118 T97620	R10606 Hs.269890	deltex (Drosophila) homolog 1 gb:yf35f11.s1 Soares fetal liver spleen
10	125120 T97775	T97775 Hs.100717	EST
	134160 T98152	T98152 Hs.79432	fibrillin 2 (congenital contractural ara
	125136 W31479	AW962364 Hs.129051	ESTs
	125144 W37999	AB037742 Hs.24336	KIAA1321 protein
15	125150 W38240	W38240	Empirically selected from AFFX single pr
	450142 W40150	AW207469 Hs.24485	chondroitin sulfate proteoglycan 6 (bama
	131987 W45435	AW453069 Hs.3657	activity-dependent neuroprotective prote
	125178 W58202	W93127 Hs.31845	ESTs
	125180 W58344	W58469 Hs.103120	ESTs
20	125182 W58650	AA451755 Hs.263560	ESTs
	446888 W68736	AL030996 Hs.16411	hypothetical protein LOC57187
	125197 W69106	AF086270 Hs.278554	heterochromatin-like protein 1
	133497 W69111	BE617303 Hs.74266	hypothetical protein MGC4251
0.5	429922 W69399	Z97630 Hs.226117	H1 histone family, member 0
25	129232 W69459	R98881 Hs.109655	sex comb on midleg (Drosophila)-like 1
	422166 W72424	W72424 Hs.112405	S100 calcium-binding protein A9 (catgran
	125209 W72724	W72724 Hs.103174	ESTs, Weakly similar to TSP2_HUMAN THROM
	125212 W72834	AA746225 Hs.103173	ESTs
30	456631 W73955	BE383436 Hs.108847	hypothetical protein MGC2749
30	125223 W74701 125225 W76540	Al916269 Hs.109057	ESTs, Weakly similar to ALU5_HUMAN ALU S
	125228 W79397	W74169 Hs.16492 AA033982 Hs.110059	DKFZP564G2022 protein
	132393 W85888	AL135094 Hs.47334	ESTs, Weakly similar to 138022 hypotheti
	125238 W86038	N99713 Hs.109514	hypothetical protein FLJ14495 ESTs
35	125247 W86881	AA694191 Hs.163914	ESTs
-	129296 W87804	Al051967 Hs.110122	ESTs
	125263 W88942	AA098878	gb:zn45g10.r1 Stratagene HeLa cell s3 93
	125266 W90022	W90022 Hs.186809	ESTs, Highly similar to LCT2_HUMAN LEUKO
	450862 W92272	U91543 Hs.25601	chromodomain helicase DNA binding protel
40	452401 W92764	NM_007115Hs.29352	turnor necrosis factor, alpha-induced pro
	428243 W93040	H05317 Hs.283549	ESTs
	125277 W93227	W93227 Hs.103245	EST
	125278 W93523	Al218439 Hs.129998	enhancer of polycomb 1
4.5	125280 W93659	Al123705 Hs.106932	ESTs
45	448205 W94003	W93949 Hs.33245	ESTs
	131844 W94401	Al419294 Hs.324342	ESTs
	125284 W94688	NM_002666Hs.103253	perilipin
	417111 W94787	AW016321 Hs.82306	destrin (actin depolymerizing factor)
50	445424 Z38294	AB028945 Hs.12696	cortactin SH3 domain-binding protein
30	125289 Z38311	T34530 Hs.4210	Homo saplens cDNA FLJ13069 fis, clone NT
	446313 Z38465 431342 Z38525	H06245 Hs.106801	ESTs, Weakly similar to PC4259 ferrifin
	433227 Z38538	AW971018 Hs.21659 AB040923 Hs.106808	ESTs kelch (Drosophila)-like 1
	428306 Z38551	AB037715 Hs.183639	hypothetical protein FLJ10210
55	424624 Z38783	AB032947 Hs.151301	Ca2+dependent activator protein for secr
	125295 Z39113	AB022317 Hs.25887	sema domain, immunoglobulin domain (ig),
	125298 Z39255	AW972542 Hs.289008	Homo sapiens cDNA: FLJ21814 fis, clone H
	125300 Z39591	Z39591 Hs.101376	EST
	448378 Z39783	BE622770 Hs.264915	Homo sapiens dDNA FLJ12908 fis, clone NT
60	444582 Z39920	R55344 Hs.22142	cytochrome b5 reductase b5R.2
	130882 Z40166	AA497044 Hs.20887	hypothetical protein FLJ10392
	128888 Z40388	AI760853 Hs.241558	ariadne (Drosophila) homolog 2
	125310 Z40646	R59161 Hs.124953	ESTs
c =	125315 Z41697	R38110 Hs.106296	ESTs
65	125317 Z99349	Z99348 Hs.112461	ESTs, Wealty similar to 138022 hypotheti
	135096 Z99394	AA081258	zinc finger protein 36 (KOX 18)

TABLE 3A

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Table 3A shows the accession numbers for those pkeys lacking unigenelD's for Table 3. The pkeys in Table 7 lacking unigenelD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession"

10	Pkey: CAT num Accession	ber: Gene d	Eos probeset identifier number tustar number ak accession numbers							
15			•							
	Pkey	CAT Number	Accession							
20	108469 124106 108501 108562 101300	116761_1 125446_1 1368412 36375_1 4669_1	AA079487 AA128547 AA128291 AA079587 AA079600 H12245 AA094769 R14576 AA083256 AA100796 AF020589 AA074629 AA075946 AA100849 AA085347 AA126309 AA079311 AA079323 AA085274 BE535511 M62098 AA306787 AW891766 AA348998 AA338869 AA344013 AW956561 AW389343 AW403607 L40391							
25 30			AW408435 AA121738 Al568978 H13317 R20373 AW948724 AW948744 AA335023 AA436722 AA448690 C21404 AW884390 AA345454 AA303292 AA174174 BE092290 T90614 AA035104 R76028 AA126924 AA741086 AW022056 AW118940 AA121666 Al832409 AA683475 Al140901 Al623576 AW519064 AW474125 Al953923 Al735349 AW150109 Al436154 AW118130 AW270782 Al804073 N27434 AA876543 AA937815 Al051166 AA505378 Al041975 Al335355 Al089540 AA662243 Al127912 Al925604 Al250880 Al366874 Al564386 Al815196 Al683526 Al435885 Al160934 H79030 Al801493 AA448691 Al673767 Al076042 Al804327 AA813438 AA680002 Al274492 T16177 Al287337 Al935050 AA907805 AA911493 Al589411 Al371358 AW576236 Al078866 AW516168 AA346372 Al560185 AA471009 R75857							
30			AA296025 AA523155 AA853168 AI696593 AI658482 AI566601 AW072797 AA128047 AA035502 AW243274 AA992517 R43760							
25	132091	94851_1	AW954243 AA829930 AA412478 AA828434 AA814538 AI927418 AI192435 W52897 AA443666 AA031913 AI683306 AA918481 AI183314 D83907 AI206832 AA876122 D83836 D83838 D82533 AI761290 AI191125 AI143749 AW771909 AI241436 AI767267 W56507 AA847787 AA568692 T10502 AI247870 AA715017 AA643304 AA890233 AA811387							
35			AA897470 AA907729 AI708679 AI078010 AA452830 AW419160 AI783713 N80205 W56778 AA676899 AI888718 N69930 AI338935 AI217580 AA639508 AA575836 BE046852 AI312651 AI038406 AA628649 AA643838 AI493761 AA032024							
•			W38849 AA340178 AA447052 AA452969 W19369 AA296364 H44229 W58767 C05751 C05835 AI741989 N98532 AW102617 AA412583 AI922246 W38495 AA355375 AA928571 C06275 AA352500 N93132							
40	117034	20113_2	U72209 NM_005748 Al655607 Al052758 AA385199 AW956794 H88679 AL135153 Al765644 AA384399 AW966458							
-1 0	100752	33207_21	AA568443 AA804610 AI873513 H88639 Z25371 R63456 W44919 T81309 BE019033 R94181 BE019198 NM_000612 J03242 AW411299 BE300064 BE297544 R94182 AW630108 T53723 D58853 H78073 H80594 BE299560 T48899 H70196 M17426 N77077 S77035 H58384 H61664 H78540 T84527 C17198 H60255 H71980 R92644 W79050 X00910 M29645 R91055 M17863 M17862 T71815 BE299561 BE464561 X06260							
45			R94741 T54216 C18594 BE262015 X06161 AW409889 AA378400 BE263228 BE313278 R88116 BE313457 H43500 T48617 BE313761 H77309 Al207601 X06159 H40413 X03425 T87663 R10627 X03562 M14118 W03982 R97520 H81229 T83157 H83168 H48762 AA669898 BE263054 H47289 AA022807 R11555 H74260 R76968 R28338 H72534 H72464 H62031 N72478 N45355 AW411300 R89113 R69135 H58454 T83281 R93476 H69645 H68015 T82229 H71089 T85121 H59939 W65299 N78176 H53099 N72373 R21788 H04660 H59639 H61874 BE262219 T53614 N73335 N50464 W00943							
50			N77189 R89257 AA570502 R89432 R06366 AA553480 AA776271 AA551359 AA551050 H51670 AA601052 BE299081 H68189 H52276 BE207832 N91192 H70332 X07868 X07868 H69464 H53782 H73710 R80435 AA553384 AW884176 N53475 T71662 AW954036 AW954033 AA552931 H93206 AA430218 AA553476 AP918470 T54124 BE207982 BE300177 N73994 AW882625 N39549 N53838 AA722389 H71878 H58909 H37849 H78435 T47933 R77174 R83814 AA411890 H94199 AA663208 BE205778 AA490137 H70492 R98232 H37800 AA679294 H40341 H74238 H47290 H73231 T48618							
55			AA025428 Al039521 H92969 N59389 H80538 H72933 T90630 AA411891 N55000 H74225 AA340290 AW957061 T54316 AA340437 H57125 H58908 H79027 H63450 N74623 R93425 H68714 H68758 N68396 H48763 N69256 H57320 H53831 H53589 N68833 N52453 H56048 H69870 H78074 R69253 R83375 T53615 H94330 H58455 H90864 T47934 H74261 R89258 R97997 R91056 R28339 R86760 H78235 R97521 H67692 H40358 AA022688 H52513 H59601 T88690 H65256 H63397 W65397 AA553588 R19280 N52645 W73930 R06367 R21743 H72372 N73921 AW883539 AW882639 T40616							
60			H47084 R95723 AA634316 AA862781 H77310 R91389 H93111 R92767 T54512 R89341 H70333 H57817 H82941 H62032 N52638 H58385 T91796 H51086 AA340292 T49918 H81230 R36121 N50411 T87664 N62436 N39340 AA665637 AA340446 H93377 H92973 BE296290 BE269788 H61665 AA340444 N54605 AA454101 R10628 R94200 Al200549 AA342640 BE298855 BE250229 T49916 H82008 N28278 AW880662 H71268 N76791 H47685 H65255 W05198 AW889144 N76677 H71702 H68036 H71915 R91612 R87807 H68059 Al133328 Al247866 AA621443 AW881050							
65			AA700847 AA340413 AW878608 AW881181 AW878249 H71916 N54596 BE161581 AW878082 W04212 AW881040 AW885492 AW880519 AA334887 AW878715 W06882 AW630222 AW885381 H70869 AW381778 H47601 AW889982 H33868 AW884986 AW878713 AW878691 AW878694 AA368070 C03393 AW878695 AW878705 AW878665 AW87878742 AW8787822 AW8787832 AW878698 R29048 AW878690 AW878686 AW878810 AW878827 AW878733 AW878659 AW878749 AW878749 AW878681 AW883353 AW883277 AW883300 AW883656 AW8783298 AW883143 AW883045							
70			AW883482 AW883352 AW883417 AW883357 AW883231 AW883474 AW883355 AW882620 AW882533 AW883754 AW883139 AW882827 AW883641 AW883567 AW883641 AW882666 AW883639 AW882827 AW882661 AW882660 AW883639 AW882830 AW882861 AW882661 AW882661 AW882630 AW882830 AW882831 AW882634 AW882630 AW882836 AW882636 AW8826 AW88							
	116417	5418_11	AW602000 AW6000000 AW602400 AW627642 AW499664 AW500888 AL042095 AW576556 AW265424 AI521500 AA761333 AA761319 AW291137 AA649040 AA769094							

AW499664 AW500888 AL042095 AW576556 AW265424 AI521500 AA761333 AA761319 AW291137 AA649040 AA769094 AA489664 AA635311 AW070509 AA425658 AI381489 AA609309 AA134476 W74704 AI923640 AW084888 H45700 AI985564 AW629495 AW614573 AI859571 AI693486 AA913892 AI806164 AA909524 AW263513 AI356361 Z40708

	123712	374423_1	Al332765 Al392620 AA181060 AW118719 AW968804 AW263502 AW505314 AA036967 W74741 R51139 H19364 H45751 Z44962 AW370823 H25650 T54007 AA453000 AL045739 AA609684 AA758732
5	117156	145392_1	W73853 AA928112 W77887 AW889237 AA148524 AI749182 AI754442 AI338392 AI253102 AI079403 AI370541 AI697341 H97538 AW188021 AI927669 W72716 AI051402 AI188071 AI335900 N21488 AW770478 W92522 AI691028 AI913512 AI144448 W73819 AA604358 N28900 W95221 AI868132 H98465 AA148793
	125008 125020	1802095_1 116017_1	T91251 T64891 T85665 T69981 T69924 AA078476
. 10	125066 116661 125104	1814993_1 1532859_1 413347_1	T86284 T81933 R61504 F04247 T95590 AA703278 H62764
	124575 125263	1666649_1 1547_2	N68168 N69188 N90450 AA098878 W88942 AW960564 AA092457 T55890 D56120 T92525 Al815987 BE182608 BE182595 AW080238 M90657 AA347236 AW961686
15	131859	3672_1	AW176446 AA304671 AW583735 T61714 AA316968 AI446615 AA343532 AA083489 AA488005 W52095 W39480 N57402 D82638 W25540 W52847 D82729 D58990 BE619182 AA315188 AA308636 AA112474 W76162 AA088544 H52265 AA301631 H80982 AA113786 BE620997 AW651691 AA343799 BE613669 BE547180 BE546656 F11933 AA376800
20			AW239185 AA376086 BE544387 BE619041 AA452515 AA001806 AA190873 AA180483 AA159546 F00242 AI940609 AI940602 AI189753 T97663 T66110 AW062896 AW062910 AW062902 AI051622 AI828930 AA102452 AI685095 AI819390 AA557597 AA383220 AI804422 AI633575 AW338147 AW603423 AW606800 AW750567 AW510672 AI250777 AA083510 AW629109 AW513200 AA921353 AI677934 AI148698 AI955858 AA173825 AA453027 AI027865 AW375542 AA454099
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25	125565	1704098_1	T91762 AA771981 Al285092 Al591386 BE392486 BE385852 AA682601 Al682884 AA345840 T85477 AA292949 AA932079 AA098791 D82607 T48574 AW752038 C06300 R20840 R20839
30	132983	11922_1	M30269 NM_002508 X82245 Al078760 AW957003 D78945 M27445 AA650439 AL048816 AV660256 AV660347 AA333052 BE295257 T60999 AA383049 AW369677 Z26985 AW175704 AA343326 AW747957 Al818389 W17308 W17302 H15591 AA371284 AA370412 W94966 BE384365 T28498 R80714 R16959 H21723 AW835154 D56097 D56381
30	118584	532052_1	W21232 AA190565 AW379755 AW067895 AW136928 AI685655 BE218584 BE465078 N68963 AA975338 BE147199 N76377 BE273749 BE397561 BE387189 AL037858 AL037878 AI963094 BE259216 AA011363 AL036189 BE562325 AA251169
35	133607	1227_6	BE617431 N98537 AA158093 AL047800 M34539 NM_000801 AA312140 D16971 AA158904 AA307114 AA312803 T09203 AW629686 AL048504 BE388578 AA220957 AA158364 BE267385 AA294971 C18055 BE241757 AA115056 Al936769 BE378435 BE206971 AW674924 BE622060 AA604674 AA115273 AW402159 AA338608 BE568819 M80199
			X55741 AA375111 AA376016 BE612671 AA805742 AW40558 N25850 N44590 H06031 AW403549 BE536552 AA056726 BE543239 AA082517 AI201645 AI201642 AI192622 N40104 AA370921 BE547569 AI969502 AA302038 AI197890 AW268354 AI014938 W45448 AI541395 AA037272 BE538826 AL039613 BE536130 AA299355 AW805147 AW974624
40			H53220 AI471471 AA399303 AA007386 W35106 BE613277 R12739 R12738 AA304342 AA687802 BE409581 AI498844 AV662092 AW904105 AA011375 BE315214 H99302 BE537893 N32299 AW855829 AI291320 BE078322 AI301395 AA303362 N32719 AA358328 AA357877 AI952540 H56279 H02758 H02048 AW805233 R82224 AA410772 AA291352
45			BE171109 N69935 BE169248 AA361173 H44978 BE617887 D52560 AA084043 W03595 R67219 N36477 N42924 R67104 H44901 H79695 W21105 AA393988 W30899 AA316096 BE622896 W46872 AA442678 BE544893 BE540112 BE621873 AA338067 N55052 BE398154 BE621210 AA740760 C03739 C03206 BE396692 AA482370 AA031614 AA301575 AA304710 AA132153 AA029796 AA994960 H19567 AA442969 H49781 H46871 AA035395 AA056185 AA149378
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50			AA587673 AI209162 AI697301 AI479995 AI679814 AI361950 AW189125 AI955888 AI986019 BE301019 AI084792 AI310211 AW189307 AI022070 AW977204 AI146825 AW190163 AW303281 AI828345 BE046043 AW029257 AA482268 AI246507 AI420729 AW084932 AW439514 AI890487 AW439692 AI523896 AI186612 AI659953 AI889773 AA687527
55			AW072694 AW262153 AW467371 Al613269 Al679238 D54404 AA158103 AW105527 AW149739 AW150361 AW268387 AW117708 Al951682 Al687440 AW674285 AA678365 Al587082 AA732095 AA019899 W45661 AA627300 BE613304 AA765891 AA612935 Al814658 AW316916 R66594 AA514640 AA025040 AA031472 AW732076 AA029797 Al244560 Al128734 AW381720 Al092360 Al263283 AW613175 Al890675 Al720156 AW631348 Al635106 Al278045 AA303979
			A1728/34 AW36172J A109236J AL263283 AW613175 A1636/5 A1720 166 AW631340 A1636 106 A12 6007 A16363 166 A17636 A AA703505 W45449 AW078661 A1292052 AW381707 A1147854 AW381743 A1518805 AA303258 AA688144 AW195967 AA428706 AA989559 AA617731 H19882 BE543418 AA830386 AA421302 W58652 T94995 A1869743 A1679145 AW085971 N98425 AA765136 A1347027 A1356955 AA928038 A1679717 AA458459 AA679281 A1367973 A1270041
60			AA765135 AA732793 AI798447 AA66646 AA251008 AI984538 AI401737 AA056186 BE043308 AW662375 AI302110 N50724 W96332 BE537047 N26983 AI567172 AA765286 AW673237 N29784 AA534275 AA084044 AW067973 AW300766 T63398 W46823 R39790 AI364185 AW298582 AA454814 AW069878 N67751 H05982 N23140 AI362647
65			AI302086 AI767772 N25755 H53114 AA706133 T93511 AA429291 AA935294 AA987647 W02803 R66595 AI680795 W23673 AW440794 AA722872 H49538 AW131042 AA531603 AA908665 AA040791 AA235312 W52205 N93444 R82180 H02759 H79696 AW088894 H56079 AA961143 AW057776 AW973745 AA016311 AW071227 AA017511 AI753994
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75			AA598917 140735 164053 AA149294 AW272948 AA365345 AAU4293 AW300697 BE261973 135001 135000 AW87876729 AW878657 AW794391 AA0699193 R01553 H44875 AA385406 AA533988 M930690 AL135600 W96331 AA017651 AA0176651 AA01849 AA017692 H85337 BE278890 AA731598 AA018512 AU076813 AU022644 R02585 X52220 AW296894 AA825671 AL9693321 AL393601 AW592611 AL146747 AA608921 AA158365 AW590007 AA354519 D20081 R02704 AW798339
, ,			M92422 AA094903 AA007676

	133681	13893_1	Al352558 Z82248 X78138 NM_003405 AU077248 AA223125 S80794 D78577 Al124697 AW403970 BE614089 BE296713
			BE621334 L20422 X80536 D54224 D54950 X57345 N29226 AA127798 AA340253 F08031 AA192540 H67636 AA321827 AW950283 AA084159 BE538808 AW401377 AA256774 C03366 W46595 W47608 AA305009 H69431 H69456 AL120082
5			H11706 AA303717 AA361357 H22042 H78020 AW999584 AA134368 AA322911 AA322961 H60980 N85248 N31547 H79624 T11718 W85826 AW894663 AW894624 BE167441 BE170015 AA304626 AW602163 AW998929 AA156681
_		•	AA151067 BE002724 AA608688 H82692 BE155392 AW383636 BE155394 AA487004 AW383504 AI342365 R82553
			W16498 BE155344 AI143938 R69901 AA322873 AW340648 R25364 AA367935 AI559406 AA033522 AA374252 AW835019 AI922133 AI697089 N99662 AW189078 AI199076 AW151598 W59944 AA662875 W94022 AA299055
10			A1039008 A1829449 AA583503 A1635674 AW131665 A1473820 AW273118 AW900930 AA908944 A1688035 AW170272
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15	114648	genbank_AA101056	AA101056
	121895	genbank_AA427396	AA427396
	100327	entrez_D55640 D55640	
	123315	714071_1 AA496369 AA	496646
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TABLE 4:

Pkey: Unique Eos probeset identifier number

Accession: Accession number used for previous patent filings

ExAcon: Exemplar Accession number, Genbank accession number

UnigenelD: Unigene number

Unigene gene title

	•		J		
10					
	Pkey	Accession	ExAccn	UniGene	UnigeneTitle
	100405	D86425	AW291587	Un 22722	nidogen 2
		D86983	D86983	Hs.118893	
15		HG1098-HT1098		Hs.121489	
		HG1103-HT1103			
		HG3342-HT3519			inhibitor of DNA binding 1, dominant neg
	100991	J03764	J03836	Hs.82085	serine (or cysteine) proteinase inhibito
	101097	L06797	BE245301	Hs.89414	chemokine (C-X-C motif), receptor 4 (fus
20		L15388	_	3Hs.211569	
	101194		L20971	Hs.188	phosphodiesterase 4B, cAMP-specific (dun
		L35545	D30857	Hs.82353	protein C receptor, endothelial (EPCR)
		L76380	_	5Hs.152175	
25		M21305 M24736	M21305 AA296520	Un 90546	gb:Human alpha şatellite and satellite 3 selectin E (endothelial adhesion molecul
23		M31166	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
		M31551	Y00630	Hs.75716	serine (or cysteine) proteinase inhibito
		M32334		Hs.347326	
		M61916	NM_002291		laminin, beta 1
30	101714	M68874	M68874	Hs.211587	phospholipase A2, group IVA (cytosolic,
		M74719		9Hs.326198	
		M92934	BE243845		connective tissue growth factor
		M94856		Hs.153179	
35		U03057		Hs.118400	
33		U03877 U18300	AA301867		EGF-containing fibulin-like extracellula damage-specific DNA binding protein 2 (4
		U27109	NM_000107	1Hs.268107	multimerin
		U31384	AW161552		quanine nucleofide binding protein 11
		U33053	U33053	Hs.2499	protein kinase C-like 1
40	102564	U59423	U59423	Hs.79067	MAD (mothers against decapentaplegic, Dr
	102663	U70322		OHs.168075	karyopherin (importin) beta 2
		U81607	NM_005100		A kinase (PRKA) anchor protein (gravin)
		U83463	AF000652		syndecan binding protein (syntenin)
45		U89942	NM_002318		lysyl oxidase-like 2
45		X04729 X06256	J03836	Hs.82085 5Hs.149609	serine (or cysteine) proteinase inhibito integrin, alpha 5 (fibronectin receptor,
		X07820	X07820	Hs.2258	matrix metalloproteinase 10 (stromelysin
		X54925	M13509	Hs.83169	matrix metalloproteinase 1 (interstitial
		X54936	BE018302		placental growth factor, vascular endoth
50		X60957	NM_005424	Hs.78824	tyrosine kinase with immunoglobulin and
		X67235		Hs.118651	hematopoietically expressed homeobox
		X67951		Hs.180909	•
		X69910	NM_006825		transmembrane protein (63kD), endoplasmi
55	103554	X79981 718951	U84722 AI878826	Hs.76206 Hs.74034	cadherin 5, type 2, VE-cadherin (vascula caveolin 1, caveolae protein, 22kD
33		AA187101		Hs.213194	
		N24990	Z44203	Hs.26418	ESTs
		R81003	AW630488		protease, serine, 23
	104764	AA025351	AJ039243	Hs.278585	
60		AA027168	AA027167		KIAA0955 protein
		AA040465	AL133035	Hs.8728	hypothetical protein DKFZp434G171
		AA045136	T79340	Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
		AA054087 AA071089	AF065214		phospholipase A2, group IVC (cytosolic,
65		AA085918	Y12059		desmoplakin (DPI, DPII) bromodomain-containing 4
03		AA187490	AA313825		AD036 protein
		AA227926	AW388633		solute carrier family 7, (cationic amino
		AA234743	AW338625		ESTs
		AA236559	AW994032	Hs.8768	hypothetical protein FLJ10849
70		AA292694	H46612		Homo sapiens HSPC285 mRNA, partial cds
		AA398243			E3 ubiquitin ligase SMURF2
		AA406363	AK001972		hypothetical protein FLJ11110
		AA411465	AB033888	MS.8519	SRY (sex determining region Y)-box 18 Homo saplens cDNA: FLJ22296 fis, clone H
75		AA412284 AA423987	X64116 H93366	Hs.7567	Homo sapiens cDNA: FLJ21962 fis, clone H
15	100124	THT LUGGI	1133300	113.1301	radio depute de la

	106155	AA425309	AA425414	Hs 33287	nuclear factor I/B
		AA435896	AA398859		hypothetical protein FLJ23221
		AA448238		Hs.16714	Rho guanine exchange factor (GEF) 15
_		AA478778	H94997	Hs.16450	ESTs
5		AA621714	BE122762		ESTs
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10		AA132983		Hs.44155	DKFZP586G1517 protein
10		AA135606			gb:zi10a05.s1 Soares_pregnant_uterus_NbH
		AA156125	AI056548		hypothetical protein FLJ20992 similar to
		AA179845		Hs.73625	RAB6 Interacting, kinesin-like (rabkines
		AA232645		Hs.42699	ESTs
15		F10399	F06838	Hs.14763	ESTs
		H16772		Hs.31444	
		N39584 N52006	AA035211 AW613287		ESTs UDP-N-acetyl-alpha-D-galactosamine:polyp
		N53375			Homer, neuronal immediate early gene, 3
20		N54067	Al287912	Hs.3628	mitogen-activated protein kinase kinase
	111133	N64436	AW580939		complement component C1q receptor
		R26892			Homo sapiens cDNA FLJ11949 fis, clone HE
		T33637	N39342		microtubule-associated protein 1B
25		T57112	H83265	Hs.8881	ESTs, Weakly similar to \$41044 chromosom
25		W80763 AA046808	AW953484	Hs.3849 Hs.108957	hypothetical protein FLJ22041 similar to
		AA253217	AI751438		40S ribosomal protein S27 isoform Homo sapiens mRNA full length insert cDN
		AA255991	AI683069		
•		AA258138		Hs.88297	ESTs
30	115819	AA426573	AA486620	Hs.41135	endomucin-2
		AA443793	R47479	Hs.94761	KIAA1691 protein
		AA490588	Al799104		Homo sapiens cDNA FLJ11333 fis, clone PL
		AA496257 AA609717		Hs.44033 Hs.66048	dipeptidyl peptidase 8 hypothetical protein FLJ10669
35		D59570	Al557212	Hs.17132	ESTs, Moderately similar to 154374 gene
	116733		AL157424		synaptojanin 2
	117023	H88157	AW070211	Hs.102415	Homo sapiens mRNA; cDNA DKFZp586N0121 (f
		H98988	H98988	Hs.42612	ESTs, Weakly similar to ALU1_HUMAN ALU S
40		N34287	AF055634		unc5 (C.elegans homolog) c
40		N52090 N66845	N52090 N66845	Hs.47420	gb:za46c11.s1 Soares fetal liver spleen
		N68905	N68905		gb:za69b09.s1 Soares_fetal_lung_NbHL19W
		R32894	BE245360	Hs.279477	
		R61715	R61715	Hs.310598	ESTs, Moderately similar to ALU1_HUMAN A
45		R71234	R71234		gb:yi54c08.s1 Soares placenta Nb2HP Homo
		R98105	C14322	Hs.250700	tryptase beta 1
	119416	W80814	T97186	He 103700	gb:ye50h09.s1 Soares fetal liver spleen Homo sapiens mRNA; cDNA DKFZp586l0324 (f
		AA404418	AA404418	115.155700	gb:zw37e02.s1 Soares_total_fetus_Nb2HF8_
50		AA405747	AW088642	Hs.97984	hypothetical protein FLJ22252 similar to
	123160	AA488687			ESTs, Wealty similar to I38022 hypotheti
		AA599143	AA599143		gb:ae52d04.s1 Stratagene lung carcinoma
		AA608588	AA608588		gb:ae54e06.s1 Stratagene lung carcinoma
55		AA608751 C13961	AA608751 C13961		gb:ae56h07.s1 Stratagene lung carcinoma gb:C13961 Clontech human aorta polyA+ mR
33		D60302		Hs.270016	
		H94892			v-ral simian leukemia viral oncogene hom
		N93521	AI680737		Homo sapiens cDNA FLJ11918 fis, clone HE
		N95477	AI571594		hypothetical protein MGC12916
60		R60044	W07701		Homo sapiens clone FLB8503 PRO2286 mRNA,
		R70506	A1887664	HS.285814	sprouty (Drosophila) homolog 4
	125103	T91518	T91518 AA570056	He 122730	gb:ye20f05.s1 Stratagene lung (937210) H ESTs, Moderately similar to KIAA1215 pro
		R45630	R60547		KIAA0372 gene product
65		R20839	R20840		gb:yg05c08.r1 Soares Infant brain 1NIB H
	125590	R23858	R23858		Homo sapiens, clone IMAGE:3840937, mRNA,
	423765		R23858		Homo sapiens, clone IMAGE:3840937, mRNA,
		AI024874	T92143	Hs.57958	
70		W26247 W26247		Hs.86859	growth factor receptor-bound protein 7 U5 snRNP-specific protein (220 kD), orth
, 0		AA856990		Hs.279531	
		AA856990		Hs.279531	
		AA136653	AW450979		gb:UI-H-BI3-ala-a-12-0-UI.s1 NCI_CGAP_Su
75		AA136653		Hs.11614	The state of the s
75		AA136653	AW450979	Un 207040	gb:UI-H-BI3-ala-a-12-0-ULs1 NCI_CGAP_Su
	12/402	AA358869	KADDOOG	115.22/949	SEC13 (S. cerevisiae)-like 1

		AI123976			MSTP031 protein
		AI123976			MSTP031 protein
		AA379500 R49693			neural proliferation, differentiation an
5		AA195678	H04150	Hs.107708	actin binding protein; macrophin (microf
,		M30257			vascular cell adhesion molecule 1
		AA028131			mesodern development candidate 1
		M10321	X06828		von Willebrand factor
	129468	J03040	AW410538		secreted protein, acidic, cysteine-rich
10	129765	M86933	M86933	Hs.1238	amelogenin (Y chromosome)
		AA012933	AA012848		tubulin-specific chaperone d
		AA286710	AF055581		lysosomal
		AA243278			mitochondrial ribosomal protein L12
15		D59711 T94452		Hs.17132 Hs.201591	ESTs, Moderately similar to I54374 gene
13		AA053400		Hs.203213	
		AA370302	D81866	Hs.21739	· · · · · · · · · · · · · · · · · · ·
		J05008	NM_00195		endothelin 1
		U85193	W27392	Hs.33287	nuclear factor I/B
20	131182	AA256153	AI824144	Hs.23912	ESTs
		X83107	F06972	Hs.27372	BMX non-receptor tyrosine kinase
		AA046593	AA040311		ESTs
		AA410480	AA359615		ESTs
25		D45304	AA443966	HS.31595	ESTs
23		M90657 AA010163	AW960564 AW361018	Hc 3383	transmembrane 4 superfamily member 1 upstream regulatory element binding prot
		AA136353	Al267615		ESTs
		Y07867		Hs.279663	
		U84573	AI752235		procollagen-lysine, 2-oxoglutarate 5-dio
30	132358	X60486	NM_003542	2Hs.46423	H4 histone family, member G
		AA132969			metalloprotease 1 (pitrilysin family)
		AA114250	AB011084		KIAA0512 gene product; ALEX2
		F13782	NM_001290		LIM domain binding 2
35		AA283035 AB002301	N92589 AB002301		ESTs, Wealty similar to I38022 hypotheti KIAA0303 protein
55		AA056731	NM_004600		Sjogren syndrome antigen A2 (60kD, ribon
		U68019	_		Homo sapiens dDNA: FLJ23037 fis, clone L
		H99198	AA125985		thymosin, beta, Identified in neuroblast
	132933	AA598702	BE263252	Hs.6101	hypothetical protein MGC3178
40	132968		AF234532		myosin X
		AA505133			done HQ0310 PRO0310p1
		AB000584	AI186431		•
		D12763 AA253193	AA026533 AW021103		interleukin 1 receptor-like 1 hypothetical protein FLJ20373
45		AA432248		Hs.183639	
		AA083572	AA403045		Homo sapiens cDNA: FLJ23197 fis, clone R
		AA479713	A1866286		ESTs, Weakly similar to B36298 proline-r
	133491	L40395	BE619053	Hs.170001	eukaryotic translation initiation factor
50	133517		NM_000165		gap junction protein, alpha 1, 43kD (con
50		W80846	Al129903	Hs.74669	vesicle-associated membrane protein 5 (m
		M34539	BE273749	NI- 75000	FK506-binding protein 1A (12kD)
	133614 133627		NM_003003 NM_002047		SEC14 (S. cerevisiae)-līke 1 glycyl-IRNA synthetase
		M85289	M85289		heparan sulfate proteoglycan 2 (perlecan
55		D10522	AI878921	Hs.75607	myristoylated alanine-rich protein kinas
		W84712	AU076964		calumenin
		D29992	C18356	Hs.295944	tissue factor pathway inhibitor 2
	133985		L34657	Hs.78146	platelet/endothelial cell adhesion motec
60		S78569	NM_002290		laminin, alpha 4
60		D43636 U97188	A1379954 AA634543	Hs.79025	KIAA0096 protein
		AA487558	AW580939		IGF-II mRNA-binding protein 3 complement component C1q receptor
		M28882	X68264		melanoma cell adhesion molecule
		X70683	Al272141	Hs.83484	SRY (sex determining region Y)-box 4
65	134656	X14787	AI750878	Hs.87409	thrombospandin 1
		AA236324	AW968058		nudix (nucleoside diphosphate linked mol
	135051		Al272141	Hs.83484	SRY (sex determining region Y)-box 4
		AA452000	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
70	135349		AA114212		serine (or cysteine) proteinase inhibito
70		D00596 D11428	X02308 NM 000304	Hs.82962 IHs 103724	thymidylate synthetase peripheral myelin protein 22 ~
		D13640			KIAA0015 gene product
		D14874	H73444	Hs.394	adrenomedullin
		D26129	NM_002933		ribonuclease, RNase A family, 1 (pancrea
75		D28476			thyroid hormone receptor interactor 12
	100405	D86425	AW291587	Hs.82733	nidogen 2

	400400	D00000	Docues	Un 440003	Molenome conscieted con-
		D86983	D86983		Melanoma associated gene
		D87953	AW888941		N-myc downstream regulated
		HG1862-HT1897			calmodulin 2 (phosphorylase kinase, delt
_		HG2614-HT2710		Hs.114599	collagen, type VIII, alpha 1
5	100619	HG2639-HT2735	N24433		RNA binding motif, single stranded inter
	100658	HG2855-HT2995	U56725	Hs.180414	heat shock 70kD protein 2
	100676	HG3044-HT3742	X02761	Hs.287820	fibronectin 1
	100718	HG3342-HT3519	BE295928	Hs.75424	inhibitor of DNA binding 1, dominant neg
	100752	HG3543-HT3739	T81309		insulin-like growth factor 2 (somatomedi
10	100828	HG4069-HT4339	AL048753	Hs.303649	
		HG417-HT417		Hs.297939	
		J03764	J03836	Hs.82085	serine (or cysteine) proteinase inhibito
		L06797	BE245301		chemokine (C-X-C motif), receptor 4 (fus
		L08246	AI439011	Hs.86386	myeloid cell leukemia sequence 1 (BCL2-r
15	101142		L12711	Hs.89643	transketolase (Wemicke-Korsakoff syndro
13					
		L13977	AA340987		prolylcarboxypeptidase (anglotensinase C
		L15388		3Hs.211569	•
	101184		NM_001674		activating transcription factor 3
20	101192		BE247295		solute carrier family 20 (phosphate tran
20		L42176	L42176	Hs.8302	four and a half LIM domains 2
	101336		NM_00673		FBJ murine osteosarcoma viral oncogene h
	101345		NM_00579		calcitonin receptor-like
	101400	M15990	M15990	Hs.194148	v-yes-1 Yamaguchi sarcoma viral oncogene
	101475	M23254	BE410405	Hs.76288	calpain 2, (m/II) large subunit
25	101485	M24736	AA296520	Hs.89546	selectin E (endothelial adhesion molecul
	101496	M26576	X12784	Hs.119129	collagen, type IV, alpha 1
		M27396	AA307680		asparagine synthetase
		M31166	M31166	Hs.2050	pentaxin-related gene, rapidly induced b
		M31994	BE293116		aldehyde dehydrogenase 1 family, member
30		M32334		Hs.347326	
-0		M35878	Al752416	Hs.77326	Insulin-like growth factor binding prote
		M36429	AF064853		guanine nucleotide binding protein (G pr
		M57730			
			NM_004428		ephrin-A1
35		M57731	AV650262		GRO2 oncogene
22		M60858	NM_005381		nucleolin
		M62994	AF043045		filamin B, beta (actin-binding protein-2
		M68874	M68874		phospholipase A2, group IVA (cytosolic,
		M69043	M69043	Hs.81328	nuclear factor of kappa light polypeptid
40		M74719			transcription factor 4
40		M75126	Al879352		hexokinase 1
		M84349	W01076		CD59 antigen p18-20 (antigen identified
	101837	M92843	M92843	Hs.343586	zinc finger protein homologous to Zfp-36
	101838	M92934	BE243845	Hs.75511	connective tissue growth factor
	101840	M93056	AA236291	Hs.183583	serine (or cysteine) proteinase inhibito
45	101857	M94856	BE550723	Hs.153179	fatty acid binding protein 5 (psoriasis-
	101864	M95787	BE392588	Hs.75777	transgelin
	101931	S76965	NM_006823	Hs.75209	protein kinase (cAMP-dependent, catalyti
	101966	S81914	X96438	Hs.76095	Immediate early response 3
	102012	U03057	BE259035	Hs.118400	singed (Drosophila)-like (sea urchin fas
50	102013	U03100	BE616287	Hs.178452	catenin (cadherin-associated protein), a
	102024	U03877	AA301867		EGF-containing fibulin-like extracellula
	102059		Al752666	Hs.76669	nicotinamide N-methyltransferase
	102121		NM_004998		myosin IE
		U31384	AW161552		guanine nucleotide binding protein 11
55		U32944	Al929721	Hs.5120	dynein, cytopiasmic, light polypeptide
		U40369	AU076887		spermidine/spermine N1-acetyltransferase
		U41767	AU077005		a disintegrin and metalloproteinase doma
		U48959			myosin, light polypeptide kinase
		U51010	U48959	HS.Z1130Z	
60			U51010	11- 70044	gb:Human nicotinamide N-methyltransferas
00		U51478	BE243877		ATPase, Na+/K+ transporting, beta 3 poly
		U53445	U53445	Hs.15432	downregulated in ovarian cancer 1
		U59289	R97457	Hs.63984	cadherin 13, H-cadherin (heart)
		U59423	U59423	Hs.79057	MAD (mothers against decapentaplegic, Dr
CE		U62015	AU076728		cysteine-rich, angiogenic inducer, 61
65		U63825	Al984144	Hs.66713	hepatitis della antigen-interacting prot
		U67963	AL119566		lysosomal
	102687	U73379	NM_007019	Hs.93002	ubiquitin carrier protein E2-C
	102693	U73824	AA532780	Hs.183684	eukaryotic translation initiation factor
	102709	U77604	AA122237	Hs.81874	microsomal glutathlone S-transferase 2
70	102759		NM_005100		A kinase (PRKA) anchor protein (gravin)
			NM_002318		lysyl oxidase-like 2
	102882		AI767736		
	102907				heme oxygenase (decyding) 1
	102915		X07820	Hs.2258	matrix metalloproteinase 10 (stromelysin
75	102927		BE512730		keratin 18
, ,					DEAD/H (Asp-Glu-Ala-Asp/His) box polypep
	102960	A 13/ 23	Al904738	Hs.76053	output (valvoirmansprins) box polypep

	103011	X52541	AJ243425	Hs 326035	early growth response 1
		X53416	X53416		filamin A, alpha (actin-binding protein-
		X54489	AW800726		GRO1 oncogene (melanoma growth stimulati
-		X54925	M13509	Hs.83169	matrix metalloproteinase 1 (interstitial
5		X57206	Y18024	Hs.78877	inositol 1,4,5-trisphosphate 3-kinase B
		X59798 X60957	AU077231 NM_005424		cyclin D1 (PRAD1: parathyroid adenomatos tyrosine kinase with immunoglobulin and
		X65965	X65965	11 13.7 0024	gb:H.sapiens SOD-2 gene for manganese su
		X69111	AL021154	Hs.76884	inhibitor of DNA binding 3, dominant neg
10		X70940	AA351647		eukaryotic translation elongation factor
		X87838			catenin (cadherin-associated protein), b
		X91247 X97748	X91247 X97748	Hs.13046	thioredoxin reductase 1 gb:H.sapiens PTX3 gene promotor region.
		Y00815	Y00815	Hs.75216	protein tyrosine phosphatase, receptor t
15		AA303711	AL120051		
		L44538		Hs.156044	
		AA025351	A1039243		
		AA027050 AA029462	AA533513 AW952619		protein disulfide isomerase related prot Homo sapiens done TCCCIA00176 mRNA sequ
20		AA045136	179340	Hs.22575	B-cell CLL/lymphoma 6, member B (zinc fi
		AA047437	AI138635	Hs.22968	Homo sapiens done IMAGE:451939, mRNA se
		AA054087	AF065214		phospholipase A2, group IVC (cytosolic,
		AA071089			desmoplakin (DPI, DPII)
25		AA156450 AA187490	AB037816 AA313825		Homo sapiens, done IMAGE:3506202, mRNA, AD036 protein
		AA195031	W84893	Hs.9305	angiotensin receptor-like 1
		AA205724	AA205759		hypothetical protein FLJ14957
		AA227926	AW388633		solute carrier family 7, (cationic amino
30		AA227986 AA234743	AA807881 AW338625		ESTs ESTs
50		AA253216	BE539071		hypothetical protein FLJ20505
		AA256210	AI805717		CGI-43 protein
		AA256268		Hs.10283	RNA binding motif protein 8B
35		AA279397 AA292379	AB024334 AL135159		tyrosine 3-monooxygenase/tryptophan 5-mo KIAA1002 protein
<i>JJ</i>		AA292717		Hs.274344	
		AA346551	AW370946		ESTs
		AA400292	W46802	Hs.81988	disabled (Drosophila) homolog 2 (mitogen
40		AA404338	AI678765	Hs.21812	ESTS
40		AA412284 AA423987	X64116 H93366	Hs.7567	Homo sapiens cDNA: FLJ22296 fis, clone H Homo sapiens cDNA: FLJ21962 fis, clone H
		AA428594	AA356392		Homo sapiens done FLB9213 PRO2474 mRNA,
		AA430108	BE019681	Hs.6019	Homo saplens dDNA: FLJ21288 fis, clone C
45		AA431462	W21493	Hs.28329	hypothetical protein FLJ14005
43		AA431470 AA443756	AL046859 AA186715	Hs.3407 Hs.336429	protein kinase (cAMP-dependent, catalyti RIKEN cDNA 9130422N19 gene
		AA449479	NM_014038		HSPC028 protein
	106634	AA459916	W25491		hypothetical protein FLJ22471
50		AA465226	N48670	Hs.28631	Homo sapiens cDNA: FLJ22141 fis, clone H
50		AA478778 AA479037	H94997 BE313412	Hs.16450	ESTs Homo sapiens done 25012 mRNA sequence
		AA482597	AF124251	Hs.26054	novel SH2-containing protein 3
		AA487561			molecule possessing ankyrin repeats indu
		AA489245	AA489245		mitogen-activated protein kinase 8 inter
55		AA504110 AA520989	AW243614		Homo sapiens cDNA FLJ10768 fis, clone NT Homo sapiens cDNA FLJ13698 fis, clone PL
		AA599434	AI817130 AL117424	Hs.9195 Hs.25035	chloride Intracellular channel 4
		AA608649	BE147611		stromal cell derived factor receptor 1
CO		AA609519	NM_012331		methionine suffoxide reductase A
60	107216		D51069	Hs.211579	
	107385	W28391	NM_005397 W28391	Hs.343258	podocalyxin-like proliferation-associated 2G4, 38kD
		AA035638	T40064	Hs.71968	Homo sapiens mRNA; cDNA DKFZp564F053 (fr
		AA083514	Al554545	Hs.68301	ESTs
65		AA121315	AB029000	Hs.70823	KIAA1077 protein
		AA147186 AA156125	AA147186 A1056548	Hs.72116	gb:zo38d01.s1 Stratagene endothelial cel hypothetical protein FLJ20992 similar to
		AA188932		Hs.132904	
		AA219653	AW007485	Hs.87125	EH-domain containing 3
70		AA232645	AW956580		ESTs .
	109737		AA055415 AW001579		ESTs, Moderately similar to A47582 B-cel
	110411 110660		AA782114		Homo saptens mRNA for KIAA1741 protein, ESTs
	110905		AA035211		ESTs
75	111018	N54067	Al287912	Hs.3628	milogen-activated protein kinase kinase
	111091	N59858	AA300067	Hs.33032	hypothetical protein DKFZp434N185

		N90933	BE301871		mannosyl (alpha-1,3-)-glycoprotein beta-
		N93764			hypothetical gene DXFZp434A1114
		R26124	AB020653		KIAA0846 protein
5		R27957 R55470	AW629414 AW083384		ESTs ESTs, Highly similar to T46395 hypotheti
,		T16550	AA307634		vacuolar protein sorting 458 (yeast homo
		T26674			Human DNA sequence from clone RP1-187J11
	113195	T57112	H83265	Hs.8881	ESTs, Weakly similar to \$41044 chromosom
	113490	T88700	BE178110	Hs.173374	Homo sapiens cDNA FLJ10500 fis, clone NT
10		T90527	H43374	Hs.7890	Homo sapiens mRNA for KIAA1671 protein,
		W42789			chromosome 8 open reading frame 4
		W60002	NM_00503		plastin 3 (T isoform)
		W78175	AA113262	HS.17901	Homo sapiens, clone IMAGE:3937015, mRNA,
15		W84768 W94427	W84768 AL035858	Un 2007	gbzh53d03.s1 Soares_fetal_liver_spleen_
13		AA253217	AI751438	Hs.41271	FXYD domain-containing ion transport reg Homo sapiens mRNA full length insert cDN
		AA426573	AA486620		endomucin-2
		AA432374	NM_00598		snail 1 (drosophila homolog), zinc finge
		AA446622	AA987568		KIAA1265 protein
20		AA478771	AJ767947		ESTs
		AA482594	D51174	Hs.272239	
		AA490588	AI799104		Homo sapiens cDNA FLJ11333 fis, clone PL
		D59570	AI557212		ESTs, Moderately similar to I54374 gene
25		H88157 H94648		Hs.293658	Homo sapiens mRNA; cDNA DKFZp586N0121 (I
2,5		H97538	W73853	10.23000	ESTs
		H98670	H45100	Hs.49753	uveal autoantigen with colled coll domai
		N22107	M18217		Homo sapiens cDNA: FLJ21409 fis, ctone C
	119559	W38197	W38197		Empirically selected from AFFX single pr
30		W80814			Homo sapiens mRNA; cDNA DKFZp58610324 (f
		AA287347	AA305599		hypothetical protein PRO2013
		AA402799	W07343 AA404418	HS.182538	phospholipid scramblase 4
		AA404418 AA425107	AI743860		gb:zw37e02.s1 Soares_total_fetus_Nb2HF8_ metallothionetn 1E (functional)
35		AA425435		Hs.300570	KIAA1204 protein
		AA442872	AL133437		Homo sapiens cDNA: FLJ21904 fis, clone H
	122577	AA452860			hypothetical protein MGC4248
		AA488687			ESTs, Weakly similar to I38022 hypotheti
40		AA599674			Homo sapiens cDNA FLJ14680 fis, clone NT
40	124059				ESTs, Weakly similar to S64054 hypotheti
		H99093 N22495	H99093		DEAD/H (Asp-Glu-Ala-Asp/His) box polypep Homo saplens mRNA; cDNA DKFZp586N0121 (f
		N23031 ·			baculoviral IAP repeat-containing 6
		R15740			carbohydrate (keratan sulfate Gal-6) sul
45		R39610	BE410405		calpain 2, (m/li) large subunit
		W45560	AL137540	Hs.102541	netrin 4
		Z39833			GTP-binding protein
		Z40583		Hs.330466	•
50		AA825437 R66613	AA825437 T40064	Hs.71968	ESTs Homo sapiens mRNA: cDNA DKFZp564F053 (fr
50		R66613	T40064	Hs.71968	Homo sapiens mRNA; cDNA DKFZp564F053 (fr
		AA868063			carbohydrate (keratan sulfate Gal-6) sul
		AA128075	AA088767	Hs.83883	transmembrane, prostate androgen induced
		AA128075	AA932794	Hs.83147	quanine nucleotide binding protein-like
55		AA128075	AA088767	Hs.83883	transmembrane, prostate androgen induced
		N66570	X69086		Homo saplens cDNA FLJ13613 fis, clone PL
		AI051390			
		AA627122 AA627122		Hs.163787 Hs.163787	
60	128453		X02761		fibroneclin 1
•		AF010193			MAD (mothers against decapentaplegic, Dr
		AA149044			type I transmembrane protein Fn14
		U82108	U82108		solute carrier family 9 (sodium/hydrogen
65		D78676		Hs.105509	
65		L35240	Z28913		enigma (LIM domain protein)
		AA598737 R69417	W28493		heat shock 70kD protein 8 STAT induced STAT inhibitor 3
		AA232837			plasmalemma vesicle essociated protein
		N72695			hypothetical protein PP1057
70		M30257			vascular cell adhesion molecule 1
		M96843	BE222494	Hs.180919	inhibitor of DNA binding 2, dominant neg
	129265	X68277	AA530892	Hs.171695	dual specificity phosphatase 1
		AA292440	R22497	Hs.110571	growth arrest and DNA-damage-inducible,
75	129468				secreted protein, acidic, cystelne-rich
75		AA228107			methionine adenosyltransferase II, beta
	101038	AA449789	DE243043	Hs.75511	connective tissue growth factor

	442724	A A A A07700	00042046	11-75544	Annua d'an la la Cartan
		AA449789	BE243845	Hs./5511 Hs.46366	connective tissue growth factor
		W01367 AA610116	AL045404		KIAA0948 protein tetraspan NET-6 protein
		AA258308	T40064	Hs.71968	Homo septens mRNA; cDNA DKFZp564F053 (fr
5		AA460273	AA453694		tripartite motif protein TRIM2
•		AA286710	AF055581	Hs.13131	lysosomal
		T68873	AA353093	16.10101	metallothlonein 1L
		D63476	D63476	Hs.172813	PAK-interacting exchange factor beta
		M62403	U20982	Hs.1516	Insulin-like growth factor-binding prote
10		X55740			5' nucleofidase (CD73)
		L10284		Hs.155560	
		AA243278	AW250380	Hs.109059	mitochondrial ribosomal protein L12
	130553	AA430032		Hs.252587	
	130638	H16402	AW021276	Hs.17121	ESTs
15		D59711	Al557212	Hs.17132	ESTs, Moderately similar to I54374 gene
	130657	T94452	AW337575	Hs.201591	
	130686	AA431571	BE548267	Hs.337986	Homo sapiens cDNA FLJ10934 fis, clone OV
		R79356	AF167706		cysteine-rich motor neuron 1
20		AA280375	AW190920		hypothetical protein SP329
20		Z49269	BE048821		small inducible cytokine subfamily A (Cy
		Z41740	A1077288	Hs.296323	•
		AA121543	AL050295		KIAA0758 protein
	131080		NM_00195		endothelin 1
25 .		AA101878		3Hs.303084	
25		T35341	AJ271216		dipeptidypeptidase III
		N87590	BE620886		GCN1 (general control of emino-acid synt
		AA256153	AI824144	Hs.23912	ESTs
		W74533 U25997	AF104266		latrophilin stanniocalcin 1
30		V01512	NM_003155 AW939251		v-fos FBJ murine osteosarcoma viral onco
50	131509		X56681	Hs.2780	jun D proto-oncogene
		AA161292	T47364		interferon, alpha-inducible protein 27
		AA491465	T93500	Hs.28792	Homo sapiens cDNA FLJ11041 fis, clone PL
		AA046593	AA040311		ESTs
35		D50914	BE559681		KIAA0124 protein
		D45304	AA443966		ESTs
		M90657	AW960564		transmembrane 4 superfamily member 1
	131909	W69127 .	NM_016558	3Hs.274411	SCAN domain-containing 1
	131915	AA316186	Al161383	Hs.34549	ESTs, Highly similar to S94541 1 clone 4
40	132046	AA384503	Al359214	Hs.179260	chromosome 14 open reading frame 4
	132050	AA136353	Al267615	Hs.38022	ESTs
		AA044755			Homo sapiens dDNA: FLJ22050 fis, done H
		U84573	Al752235	Hs.41270	procollagen-lysine, 2-oxoglutarate 5-dio
15		AA058911	AA235709		DKFZP586O1624 protein
45		AA620962		Hs.325093	
		AA285290 X60486		Hs.323806	pinin, desmosome associated protein
		R31641	NM_003542 AA876616		H4 histone family, member G ESTs, Weakly similar to A43932 mucin 2 p
		AA489190	AW163483		double ring-finger protein, Dorlin
50	132490		NM_001290		LIM domain binding 2
50		AA257993	AA257992		Janus kinase 1 (a protein tyrosine kinas
		M24283	M24283		Intercellular adhesion molecula 1 (CD54)
		AA443114	AA160511		amino acid system N transporter 2; porcu
	132716		BE379595	Hs.283738	caseln kinase 1, alpha 1
55	132840	N23817	BE218319		GTPase Rab14
	132883	AA047151	AA373314	Hs.5897	Homo sapiens mRNA; cDNA DKFZp586P1622 (f
	132958	N77151	AF234532		myosin X
		AA480074			
60		Y00787	Y00787	Hs.624	Interleukin 8
60		T99789	BE384932		ESTs, Weakly similar to AF257182 1 G-pro
		W84341	AW946276		Homo saplens mRNA; cDNA DKFZp586J021 (fr
		L09209	W16518		amytoid beta (A4) precursor-like protein
		D12763	AA026533		interleukin 1 receptor-like 1
65	133149		AA370045 AW021103		AXIN1 up-regulated
05		AA253193 AA432248	_		hypothetical protein FLJ20373
	133220			Hs.183639	hypothetical protein FLJ10210 Homo sapiens mRNA full length insert cDN
		AA083572	AA403045		Homo sapiens cDNA: FLJ23197 fis, done R
	133295		AJ147861		tow density lipoprotein receptor (famili
70		N75791	AW631255		L-3-hydroxyacyl-Coenzyme A dehydrogenase
		X57579	AW103364		inhibin, beta A (activin A, activin AB a
		X02612	NM_000499		cytochrome P450, subfamily I (aromatic c
		H44631	BE294068		Immediate early protein
		AA090257		Hs.177781	
75	133478		X83703	Hs.31432	cardiac ankyrin repeat protein
	133491				eukaryotic translation initiation factor
			-		·

	133510	AA227913	AW880841	Hs.96908	p53-induced protein
		X52947	NM_00016		gap junction protein, alpha 1, 43kD (con
		M11313	AU077051		alpha-2-macroglobulin
		L14837	NM_00325		fight junction protein 1 (zona occludens
5		M60721	M60721	Hs.74870	H2.0 (Drosophila)-like homeo box 1
_		D90209	D90209		activating transcription factor 4 (tax-r
		T67986	T70956	Hs.75106	clusterin (complement lysis inhibitor, S
		AA148318		Hs.75249	ADP-ribosylation factor-like 6 interacti
		U97105	Al301740	Hs.173381	dihydropyrimidinase-like 2
10	133671	T25747	AW503116	Hs.301819	
	133678	K02574	AW247252		nucleoside phosphorylase
	133681	D78577	Al352558		tyrosine 3-monooxygenase/tryptophan 5-mo
	133722	X53331	AW969976	Hs.279009	
	133730	S73591	BE242779	Hs.179526	upregulated by 1,25-dihydroxyvitamin D-3
15	133750	X95735	BE410769	Hs.75873	zyxin
•	133802	L16862	AW239400	Hs.76297	G protein-coupled receptor kinase 6
	133825	U44975	BE616902	Hs.285313	core promoter element binding protein
		M97796	BE222494	Hs.180919	Inhibitor of DNA binding 2, dominant neg
		U86782	U86782	Hs.178761	26S proteasome-associated pad1 homolog
20		AA099391	U48959	Hs.211582	myosin, light polypeptide kinase
		M19267	M19267	Hs.77899	tropomyosin 1 (alpha)
		D29992	C18356	Hs.295944	
		L19314	Al125639		
0.5		S78569	_	DHs.78672	laminin, alpha 4
25		U28811	NM_01220		Golgi apparatus protein 1
		L77886	AL034349		protein tyrosine phosphatase, receptor t
		C14407	AW245540		brain abundant, membrane attached signal
		M60278	AA161219		diphtheria toxin receptor (heparin-bindi
20		R81509		Hs.160726	
30		AA487558	AW580939		complement component C1q receptor
		D86962	D86962	Hs.81875	growth factor receptor-bound protein 10
		AA478971	R70429	Hs.81988	disabled (Drosophila) homolog 2 (mitogen
		D50683	D50683	Hs.82028	transforming growth factor, beta recepto
35		U56637	AI557280	MS.1842/U	capping protein (actin filament) muscle
22		M61199	AA334551	U= 244570	sperm specific antigen 2
		M28882 X15183	X68264		melanoma cell adhesion molecule
		S53911	M30627	75.269066 3Hs.85289	heat shock 90kD protein 1, alpha
		U20734			CD34 antigen jun B proto-oncogene
40		D28235	D28235		prostaglandin-endoperoxide synthase 2 (p
40		AA236324	AW968058		nudix (nucleoside diphosphate linked moi
		AA148923	AL136653		decidual protein induced by progesterone
		AA174183			KIAA1682 protein
		AA456311	AA876372		Homo sapiens mRNA; cDNA DKFZp667D095 (fr
45		L08069	W27190	Hs.94	DnaJ (Hsp40) homolog, subfamily A, membe
		AA452000	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f
		AA282140	T53169	Hs.9587	Homo sapiens cDNA: FLJ22290 fis, clone H
		J02854	C03577	Hs.9615	myosin regulatory light chain 2, smooth
		AA442054	U80983	Hs.268177	

TABLE 4A

Table 4A shows the accession numbers for those pkeys tacking unigenelD's for Table 4. The pkeys in Table 7 tacking unigenelD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligoniudeotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" 5

10 Pkey: CAT number: Unique Eos probeset identifier number

Accession:

Gene cluster number Genbank accession numbers

15	Pkey	CAT Number	Accession
	100752	33207_21	T81309 BE019033 R94181 BE019198 NM_000612 J03242 AW411299 BE300064 BE297544 R94182 AW630108 T53723 D58853 H78073 H80594 BE299560 T48899 H70196 M17426 N77077 S77035 H58384 H61664 H78540 T84527 C17198 H20256 B74089 H70196 M17426 N77077 S77035 H58384 H61664 H78540 T84527 C171198 H20256 B74089 H70196 M17426 M17893 H70197 H7019
20			H60255 H71980 R92644 W79050 X00910 M29645 R91055 M17863 M17862 T71815 BE299561 BE464561 X06260 R94741 T54216 C18594 BE262015 X06161 AW409889 AA378400 BE263228 BE313278 R88116 BE313457 H43500 T48617 BE313761 H77309 AI207601 X06159 H40413 X03425 T87663 R10627 X03562 M14118 W03982 R97520 H81229 T83157 H83168 H48762 AA669898 BE263054 H47289 AA022807 R11555 H74260 R6745 B6864 B6874 R3281 R93476 H69664 51680 T83229 H71089 T85121 H50320 W65200 N76475 M52000 N776478 M52000 N
25			H59939 W65299 N78176 H53909 N72373 R21788 H04660 H59639 H61874 BE262219 T53614 N73335 N50464 W00943 N77189 R89257 AA570502 R89432 R06366 AA553480 AA776271 AA551359 AA551050 H51670 AA601052 BE299081 H68198 H52276 BE207832 N91192 H70332 X07868 K07868 H69464 H53782 H73710 R80435 AA553384 AW884176 N53475 T71662 AW954036 AW954033 AA552931 H93206 AA430218 AA553476 A1918470 T54124 BE207982 BE300177 N73994 AW882625 N39549 N53838 AA722389 H71878 H58909 H37849 H78435 T47933 R77174 R83814 AA411890 H94199 AA663208 BE205778 AA490137 H70492 R98232 H37800 AA679294 H40341 H74238 H47290 H73231 T48618
30			AA025428 Al039521 H92969 N59389 H80538 H72933 T90630 AA411891 N55000 H74225 AA340290 AW957061 T54316 AA340437 H57125 H58908 H79027 H63450 N74623 R93425 H68714 H68758 N68396 H48763 N69256 H57320 H53831 H53589 N68833 N52453 H56048 H69870 H78074 R69253 R83375 T53615 H94330 H58455 H90864 T47934 H74261 R89258 R97997 R91056 R28339 R86760 H78235 R97521 H67692 H40358 AA022688 H52513 H59601 T88690 H65256
35			H63397 W65397 AA553588 R19280 N52645 W73930 R06367 R21743 H72372 N73921 AW883539 AW882639 T40616 H47084 R95723 AA634316 AA862781 H77310 R91389 H93111 R92767 T54512 R89341 H70333 H57817 H82941 H62032 N52638 H58385 T91796 H51086 AA340292 T49918 H81230 R36121 N50411 T87664 N62436 N39340 AA665637 AA340446 H93377 H92973 BE296290 BE269788 H61665 AA340444 N54605 AA454101 R10628 R94200 AI200549 AA342640 B93377 H92973 BE296290 H24916 H82008 N28278 AW880662 H71268 N76791 H77685 H65255 W05198
40			AW889144 N76677 H71702 H68036 H71915 R91612 R87807 H68059 Al133328 Al247866 AA621443 AW881050 AA700847 AA340413 AW878608 AW881181 AW878249 H71916 N54596 BE161581 AW878082 W04212 AW881040 AW885492 AW880519 AA334887 AW878715 W06882 AW630222 AW885381 H70869 AW381778 H47601 AW889982 H63868 AW884986 AW878713 AW878685 R36391 AW878694 AA368070 C03393 AW878695 AW878705 AW878665 AW878742 AW878620 AW878623 AW878688 R29048 AW878690 AW878686 AW878810 AW878827 AW87873
45			AW878659 AW878749 AW878681 AW883353 AW883277 AW883300 AW883555 AW883228 AW883143 AW883045 AW883482 AW883352 AW883417 AW883357 AW883231 AW883355 AW88355 AW882533 AW883754 AW883139 AW882827 AW883641 AW883567 AW883481 AW882983 AW882982 AW882465 AW883419 AW882466 AW883639 AW882230 AW882981 AW882534 AW882874 AW882619 AW883480 AW882826 AW882831 AW882835 AW882830 AW883563 AW882456 AW627642
50	117156	145392_1	W73853 AA928112 W77887 AW889237 AA148524 AI749182 AI754442 AI338392 AI253102 AI079403 AI370541 AI697341 H97538 AW188021 AI927669 W72716 AI051402 AI188071 AI335900 N21488 AW770478 W92522 AI691028 AI913512 AI144448 W73819 AA604358 N28900 W95221 AI888132 H98465 AA148793
55	131859	3672_1	AW960564 AA092457 T55890 D56120 T92525 Al815987 BE182608 BE182595 AW080238 M90657 AA347236 AW961686 AW176446 AA304671 AW583735 T61714 AA316968 Al446615 AA343532 AA083489 AA488005 W52095 W39480 N57402 D82638 W25540 W52847 D82729 D58990 BE619182 AA315188 AA308636 AA112474 W76162 AA088544 H52265 AA301631 H80982 AA113786 BE620997 AW651691 AA343799 BE613669 BE547180 BE546656 F11933 AA376800
			AW239185 AA376086 BE544387 BE619041 AA452515 AA001806 AA190873 AA180483 AA159546 F00242 Al940609 Al940602 Al189753 T97663 T66110 AW062896 AW062910 AW062902 Al051622 Al828930 AA102452 Al685095 Al819390 AA57597 AA383220 Al804422 Al633675 AW338147 AW603423 AW606800 AW750567 AW510672 Al250777 AA083510
60			AW629109 AW513200 AA921353 AI677934 AI148698 AI955858 AA173825 AA453027 AI027865 AW375542 AA454099 AA733014 AI591384 R79300 R80023 AA843108 AA626058 AA844898 AW375550 AA889018 AI474275 AW205937 AI052270 AW388117 AW388111 AA699452 AI242230 N47476 H38178 AA366621 AA113196 AA130023 H39740 T61629 AI885973 AW083671 AA179730 AA305757 AI285455 N83956 AA216013 AA336155 AW999959 T97525 AA345349 T91762 AA771981 AI285092 AI591386 BE382486 BE385852 AA682601 AI682884 AA345840 T85477 AA292949
65	125565 133607	1704098_1 1227_6	AA932079 AA098791 D82607 T48574 AW752038 C06300 R20840 R20839 BE273749 BE397561 BE387189 AL037858 AL037878 AI963094 BE259216 AA011363 AL036189 BE562325 AA251169 BE617431 N98537 AA158093 AL047800 M34539 NIM_000801 AA312140 D16971 AA158904 AA307114 AA312803 T09203 AW629686 AL048504 BE388578 AA220957 AA158364 BE267385 AA294971 C18055 BE241757 AA115056
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75			AV662092 AW904105 AA011375 BE315214 H99302 BE537893 N32299 AW855829 Al291320 BE078322 Al301395 AA303362 N32719 AA358328 AA357877 Al952540 H56279 H02758 H02048 AW805233 R82224 AA410772 AA291352

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5			BE171109 N69935 BE169248 AA361173 H44978 BE617887 D52560 AA084043 W03595 R67219 N36477 N42924 R67104 H44901 H79695 W21106 AA393988 W30899 AA316096 BE622896 W46872 AA442678 BE544893 BE540112 BE621873 AA338067 N55052 BE398154 BE621210 AA740760 C03739 C03206 BE396692 AA482370 AA031614 AA301575 AA304710 AA132153 AA029796 AA994960 H19567 AA442969 H49781 H46871 AA035395 AA056185 AA149378 AA643080 AL135479 AA292329 AA654337 AA041228 AA454888 AA025039 W58331 AA625981 T94941 AA302448 H19900 AA218956 AA513790 AA563962 AA398076 W44441 AA293276 W47373 AA625879 W30688 AA043029 T64284
10			R79151 AA304340 AA485186 AA604939 R82470 AA421425 AW771456 AI339329 AA304424 AA605236 AA936934 AA587673 AI209162 AI697301 AI479995 AI679814 AI361950 AW189125 AI955888 AI986019 BE301019 AI084792 AI310211 AW189307 AI022070 AW977204 AI146825 AW190163 AW303281 AI828345 BE046043 AW029257 AA482268 AI246507 AI420729 AW084932 AW439314 AI890487 AW439692 AI523896 AI186612 AI659953 AI659953 AI659953 AI659612 AI668073 AI6687527
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60		·	H01101 H01867 AA548743 Al671806 AW872949 AW872941 AA742447 Al199788 AA045604 Al637465 Al741796 AW242217 AW131463 Al765302 Al683923 AA889762 Al804889 Al986437 C06049 BE502340 Al695651 Al491970 AA496804 AA281008 AA665699 Al473814 BE301445 AA707837 AA551925 Al017348 Al208185 AA775203 AA156296 AA557463 H95441 AA768547 AW769358 AA991197 AA181954 Al091389 Al147289 AW771837 Al638582 AA844411 Al374750 T29320 AW951272 AW085923 H02834 AA843259 AA814696 AW183290 AA158453 N68125 N69039 AA100423
65	126872	142696_1	AA101346 AI918720 H01102 R67868 H01868 N66438 R46580 AI858433 AA599560 AA187577 AA157481 AA361520 AL047827 AA158452 R21688 AW964874 AA325161 R40871 AW752395 AW375924 R13355 AA281174 AA428908 AW450979 AA136653 AA136656 AW419381 AA984358 AA492073 BE168945 AA809054 AW238038 BE011212 BE011359 BE011367 BE011368 BE011362 BE011365 BE011363
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75	123523	genbank_AA608	AA639462 Al261373 Al432414 Al984994 Al539335 AA401550 AA358757 Al609976 AA442357 AA359393 AA437046 AA370301 AA429328 AW272055 Al580502 Al832944 Al038530 AA425107 Al014986 Al148349 AW237721 AW779756 AW137877 Al125293 AA400404 R28554 8588 AA608588

	123533 125091	genbank_AA608751 AA608751 genbank_T91518 T91518
	123964 102491	genbank_C13961 C13961 entrez_U51010 U51010
5	118475 118581 113947 101447	genbank, N66845 N66845 genbank, N68905 N68905 genbank_W84768 W84768 entrez_M21305 M21305
10	101667	13349_1 NM_005381 M60858 AW373732 AW373724 AW373689 AW373629 AW373609 AW373776 AA187806 AW386946 AW374207 T05235 AA216203 AW386556 AA306940 AA306526 AA315461 AL036757 AW373711 AW403124 AW403640 AW377084 T27360 H62638 F06957 AW377051 AA554779 AA378568 AA096007 AW352407 AW302637 F07929 H17433 AW382712 H05665 F07292 N39875 AA089729 H62556 N42842 R12952 AW3737355 AW364155 AA056183 W39185 AW382708 N32488 AF114096 AW375993 Al133569 W52561 AA603040 AA133710 Al928796 AW176370 AA827519
15		AW338437 AA521142 T29341 AI800461 AW317002 AA703914 AA860830 AI859203 AI445772 AA714334 AI817066 AI832027 AW510442 AI635802 AW08806 AW068572 AW408555 AW467542 AA552657 AA152367 W32081 AA582124 AA074040 AA931657 AI051154 AW410203 AI921644 H17434 AI832330 AW404836 AI925038 AA088423 AA954166 AA580453 AW021292 AI267215 AW080082 AW383778 AI933053 AI919097 W31557 N90245 AA931591 AA563995 F36352 AA056184 AA476294 AA641327 AA533550 AI749630 W58323 AA569119 AA508573 AI809050 AI378996
20		AA411362 AW407505 AA938104 AA074041 AA632876 AW193748 AA507873 Al270128 Al472365 AA411363 Al523216 AJ719965 Al816302 AA182681 AJ707990 AA133588 AJ758537 W60253 Al460308 AA135423 Al083904 F04188 N89693 AW408776 ALG78595 Al270568 AA722059 W58234 F33650 AA090547 AA285108 AA425981 N85079 D20218 Al273980 AA159028 F03226 AW247914 N26918 AW272741 N90109 H05666 N23327 AW247953 R44748 AA962015 F03558 AJ752394 AW409913 AW248396 Al816463 AJ752393 AA325370 AA263089 Al570130 Al971951 Al160658 AJ357360
25 -		AW168686 AL.121075 AW050536 N21672 W67748 AA514242 AI127386 H14607 AI185752 W79364 AA088520 AA152476 AW351940 AW373683 AI940524 AW374953 T56500 N24329 AI940720 AW374933 AW374947 AW391913 AL.138337 AW376241 AW062943 F26666 AW410202 AW062958 F34529 AW381807 AW383315 W17147 AW176359 AA664576 AW380424 AA306040 AI745674 AW300951 AI188579 AI438973 AI305271 AA433818 AA612807 AI831809 AI940409 AA155663 AI572988
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	103432 119174 133678	entrez_X97748 X97748 genbank_R71234 R71234 11235_1 AW247252 AA346143 NM_000270 AA381085 N91995 X00737 AA381079 AA296473 AA296110 AA315735 AA311617
35		AA326750 AA376804 AW403290 T95231 M13953 T47963 H82039 AA279899 AA627997 N76320 N99527 H37842 W20095 AA457308 AW469547 AA724143 H83220 AA319496 W86334 W30892 R89169 R99427 N41854 H47286 AA348094 AA045089 R63016 AI922219 AI024906 AI096488 AI885005 AA194872 N90489 AI452544 H72411 AA282427 AA430735 R68963 R22453 H70385 AW129369 AW467320 AW519082 AA345018 AA582183 AI961789 R65918 N30611 AI979189 AI280889 AW273191 R66531 AI285845 AI675927 AI421990 AW190879 H37794 AA699667 H68427 AA954388
40		A1188757 A1140048 AA430382 A1204151 AW247864 AA559099 AI431420 AA548276 A1149466 AA772669 AA694388 AA724168 AA301651 AA281952 AA779925 AA234760 W86290 AA913603 AW511745 AI500697 AA814922 AA635040 T47964 H53998 AA975804 R98710 A1077604 N70252 R98084 AW250171 H69268 AI597614 AA970746 AA972548 A1377116 R62962 H16737 R89070 AA731329 R66532 N54364 AI818832 H81944 N71567 T95122 W86463 AA437095 A1431999 AI915724 N63851 AI674743 AA457307 AA211475 N64444 A1799146 H72853 R99335 H60413 AA770367
45		AA156105 Al269937 H64029 H89728 R65819 AW470496 Al873318 Al735713 H82987 C02447 Al478666 T27651 Al699770 AW025156 H69719 Al984717 N69225 Al459856 AA953577 Al424691 H13843 R22404 Al873796 Al336002 N70898 Al420854 AA541792 AA346142 Al000814 Al828348 AA045090 T51257 N90434 H13890 N73184 Al708083 AA781606 AA329050 AA339985 R68964 H64795 W04186 H16845
50	119416 119559	genbank_T97186 T97186 NOT_FOUND_entrez_W38197 W38197
JU	123473	genbank_AA599143 AA599143 .

TABLE 5:

Unique Eos probeset identifier number Accession number used for previous patent filings Exemplar Accession number, Genbank accession number Unigene number Unigene gene title Pkey: Accession: ExAccn: UnigeneiD: Unigene Title: 5

10					
	Pkey	Accession	ExAccn	UniGene	UnigeneTitle
15	115819 132837 101545 102898	AA426573 D58024 M31210 X06256	AA486620 AA370362 BE246154 NM_002205	Hs.57958 Hs.154210	AA486620 AA370362 BE246154 NM_002205
20	105330	L20859 X07820 AA234743 U97519 U03877	BE247295 X07820 AW338625 NM_005397 AA301867	Hs.2258 Hs.22120 Hs.16426	BE247295 X07820 AW338625 NM_005397 AA301867
25	104865 106124	M28882 X54925 AA045136 AA423987 AA234743	X68264 M13509 T79340 H93366 AW338625	Hs.211579 Hs.83169 Hs.22575 Hs.7567 Hs.22120	X68264 M13509 T79340 H93366 AW338625
30	109001 104764 133200 105263 105178 109456	AA156125 AA025351 AA432248 AA227926 AA187490 AA232645	AA313825	Hs.72116 Hs.278585 Hs.183639 Hs.6682 Hs.21941 Hs.42699	Al056548 Al039243 AB037715 AW388633 AA313825 AW956580

TABLE 5A

Table 5A shows the accession numbers for those pkeys tacking unigenelD's for Table 5. The pkeys in Table 7 tacking unigenelD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTvist, Oaldand California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10 Pkey: CAT number:

5

Unique Eos probeset identifier number Gene cluster number Genbank accession numbers

Accession:

15	Pkey	CAT Number	Accession
20	115819	10241_1	AA486620 AF205940 AA297524 AB034695 AA081335 NM_016242 AA188323 AA297537 H88204 AW953081 W31695 AW582203 AA248250 AW681211 AA426230 AA464807 AA426155 N44141 AA347390 AA770661 AI333225 N36136 AW665724 AA431894 AI374976 AI400254 AI338446 AA186695 H88205 W04527 AA487056 AI051414 AA918383 AA426573 AA425620 AW433654 AA090513 BE167284 BE167291 AI301726
20	102024	14505_1	AA301867 AW957981 R27614 AA155808 AI920990 AI740711 AA301026 AA301015 AI220981 AI857670 AI537140 AW015210 AA030000 W46890 H44021 AI355967 AI651735 AA058479 AA146932 T58265 R85890 AA047810 AA017387 AW026093 AA971133 AI827263 AI056416 AI355994 AI127691 H46603 U03877 NM_004105 AA157357 H42844
25			AA146824 AA187709 AA187269 AA304348 AA147292 AA361687 AA156041 AA330636 R32929 AA321130 AW950260 AA082157 AA029129 AA303708 AA028155 D31561 T84689 AA302493 BE153057 BE153181 W39408 AA187200 BE153250 AW383337 AW382622 AW382647 AW750072 BE153060 AW382630 AW371865 AW392464 AW382664 AW382658 AW382650 H61647 AW365075 AW365049 AA373397 BE072779 BE072781 Z30254 W24381 BE153254 AA040442 BE072779 BE072731 N94740 AA146945 AW802737 AI826799 AI085395 R34034 H65140 AA082800 H88275
30			AA147824 R63882 W80899 AA296413 AI765300 AI862426 AW022055 AW300003 AI743784 AI862635 AI985428 AA147764 AW573245 AW190290 AI040898 D57613 N63457 AA148082 AI028458 AA148110 AW814489 N75105 AW629443 AA704122 AW582220 AA181240 AA057495 AI418224 AI261751 AW388595 AI472205 AW470672 AA102546 AA789046 AA182416 AA062668 AW300732 AI288220 AA181982 AA146825 AA028130 AI985522 AA303344 AA081313 N69082 AA182035 AI867128 AA100902 AA605087 N67178 AW020324 AW890446 AI472191 AI335691 AI597837
35			AI081143 AI335681 AA040443 AI128067 AI678244 AA018303 AA157260 W80792 AI934590 AI096430 T54343 AI446350 AA165196 AA780683 AA603631 AA047787 AA968580 AA912645 AW890504 AW026913 D56983 H52088 AA156121 R30848 AW023036 AI5990960 N67345 AI753225 AI753228 AI83768 AH7818 H89101 AI362141 H89205 AI147711 AA321129 AA668622 AA343479 AW069438 AI422376 AW629270 AA013413 AI221948 AA970605 N52335 H38366 T91180 AA657841 AA017386 AA152227 AA187593 AI913340 AI719313 AI969943 AI701271 AI004328 AI868348 N93659 H65093 H25736 D57007 D56957 C00987 D61839 D56661 AI472137 AI971002 D56971 BE048830 D57972 AI589286
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45	109456	180633_1	AW956580 AA886361 Al147670 Al090115 Al168683 AA232645 H99504 AA374707 AA380875 AW139567 Al735132 BE439385 AW629780 N28322 AA232789 AA232790 N73285
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55	133200	28960_1	Al613182 N94510 W47343 Al085755 Al076956 Al918426 AA081208 Al282835 AA147528 Al081490 Al654536 AA181875 AA081282 AA186389 C06085 AA083542 Al800644 AA157642 AA101069 AA157752 AA158121 AA143331 AA081283 AA852603 AA188296 Al932880 AW449628 AA187348 C02091 AA514656 AA082736 AA308786 AA143201 M16567 AB037715 Al351347 Al375796 Al884765 AL121124 W01068 Al807275 T95240 R42807 AW515645 Al057314 Al033520
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65			AW022003 Al862704 H19344 R61511 Al080204 H16566 AA432248 Al767980 T16688 Al984342 Al217478 Al767095 Z38551 Al359566 Al361437 Al041000 R07033 H16608 H19054 R12874 R61567 N98368 BE221199 Z42320 AA094554 R07078 AW850886 AA418090 R41262
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, ,	102898	24023_1	NM_002205 X06256 M13918 BE070866 AW239485 AW996127 BE273894 BE272590 BE410252 R25975 T11786 T11787 AA301142 AA301165 AW960506 BE272819 AA386086 T39391 AA285303 AA370580 D58585 T58668 AA156213 W24142 AA343323 AW796067 AA151197 AA376121 R94782 AA302363 H90357 R82621 AA301677 H55997 AW796059 W92358
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	134416	30694_1	X68264 NIM_006500 AF089868 BE257461 BE275425 AW997154 AI902799 AI902803 M78206 AA085691 AW392972
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			TOOLOG TOO TOO TOO TOO TOO TOO TOO TOO TOO T

AV653575 R27900 N48215 AW366371 N45500 AV652967 AI889251 Al080457 N39021 AI738542 AW242849 Al857471
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Al393465 AW069210 AI743830 AA744243 AA401310 AW439758 AW088152 R93391 AA291379 AA225220 AW009358
Al192879 AA291202 Al565089 AA225089 AA807688 Al052058 Al341641 Al066625 AA333864 AA159147 Al923912
R75851 AI761143 AW768588 AA394195 Al288450 AW512564 Al452775 Al056520 AA468602 AA872566 Al434739
AA291838 Al948623 AW768614 Al374753 AW068174 AA884908 Al199347 W94946 Al159995 AA877642
Al280646 Al307610 AA403310 R08205 AW182123 Al000999 R27808 AW026571 D20816 Al560350 T27667 AW960271
Al174628 Al432042 Al424528 AA909562 T17342 Al783866
Al056548 AW409843 AW263540 AA723669 AA909334 AA156120 AA157141 AA156125 AW409866 W19499 AA157229
AW887435

TABLE 6:

5

Pkey: Unique Eos probeset identifier number

ExAccn: Exemplar Accession number, Genbank accession number

UnigenelD: Unigene number

Unigene Title: Unigene gene title

AUC1: 70th percentile of average intensity (AI) for probeset at each of 2,6,15,24,48, and 96 hour timepoints minus 70th percentile AI at 0 hrs, summed over 5 experiments.

AUC2: AUC1/90th percentile of AI for aorta, aortic valve, vein, and artery.

10

	Pkey	Ex.Accn	UnigenelD	UnigeneTitle	AUC1	AUC2
15	314941	AA515902	Hs.130650	ESTs	1038	9
	327414			predicted exon	303.2	30.3
	321911	AF026944	Hs.293797	ESTs	429.2	42.9
	331578	A1246482	Hs.249989	ESTs	677.4	10.3
	332466	AB018259	Hs.118140	KIAA0716 gene product	395.2	39.5
20	313513	AW298600	Hs.141840	ESTs, Weakly similar to S59501 interfero	324	32.4
	320635	N50617	Hs.80506	small nuclear ribonucleoprotein polypept	394.8	39.5
	326230			predicted exon	357.2	35.7
	313556	AA628517	Hs.118502	•	433.6	12
	313665	AW751201	Hs.120932	ESTs	-83	0.5
25	324852	Al380792	Hs.135104	ESTs	348.2	34.8
	314372	AL040178	Hs.142003	ESTs, Weakly similar to The KIAA0149 gen	-49.2	0.5
	311877	AA084248	Hs.85339	G protein-coupled receptor 39	-1309	0.2
	322262	AA632012			-247.8	1
	312173	AI821409	Hs.304471	ESTs, Highly similar to AF116865 1 hedge	-1025.8	1
30	319795	AB037821	Hs.146858	protocadherin 10	203.6	5.2
	313350	AW591949	Hs.57958	ETL protein	183.8	18.4
	326759			predicted exon	1654.4	1.2
	300318	AW444502	Hs.256982	ESTs, Highly similar to AF116865 1 hedge	-346	1
	313978	A1870175	Hs.13957	ESTs	576.6	2.3
35		A1077477			56.4	0.4
				semaphorin Rs, short form	-127.6	0
		BE547674			-102.6	0
	321325	AB033100	Hs.300646	KIAA protein (similar to moușe paladin)	1080.6	4.8
40	303251	AF240635	Hs.115897	protocadherin 12	1270.8	5.3
40				Homo sapiens mRNA full length insert cDN	915.8	15.8
				ESTs, Moderately similar to ALUC_HUMAN I		4.9
		AW337575	Hs.201591		522.6	4.7
	337214			predicted exon	269	26.9
4.5		AW023595	Hs.232048		796.4	20.2
45		AA582081		gb:nn32h08.s1 NCI_CGAP_Gas1 Homo sapk		316.4 10.5
		AA644669			349.6	7.6
		AW192334			638.6	63.9
		AA972965			360.8	36.1
50				RNA binding motif protein, X chromosome	700.2	6.6
50		T90309	Hs.269651		274.2	7.5
	319267		Hs.6818	ESTS	238.2	23.8
		H75391	Hs.255748		231.8	23.2 8.2
	326198	USEDOO	U- 004F04	predicted exon	581.6	
55		H25899	Hs.201591		281.6	9.7
22		AW072215			-213 285	0.3
		W87874	Hs.25277	hypothetical protein FLJ21065	26.6	0.5 0.3
	_	DE301700	HS.233933	hypothetical protein FLJ20401	1494.2	34.7
	338316	R44557	Hs.23748	predicted exon ESTs	975.8	1.8
60		NM_006033		lipase, endothelial	201.2	0.9
00				Homo sapiens mRNA; cDNA DKFZp434B042		478.6 1.3
		X17033		Integrin, alpha 2 (CD49B, alpha 2 subuni	356.2	1.7
	325544	A17000	115.27 1300	predicted exon	1014.6	9.4
	328700			predicted exon	627.4	62.7
65		AW248508	He 279727	Homo sapiens cDNA FLJ14035 fis, clone HE		5.7
05	336034	7111210000	113.213121	predicted exon	782.6	78.3
		AA938198	He 146123	hypothetical protein FLJ12972	746.4	13.8
		AW341683	120120	gbthd13d01x1 Soares_NFL_T_GBC_S1 Hon		134.8 13.5
		R39288	Hs.6702	ESTs	137	13.7
70		H06350		Human DNA sequence from clone RP5-850E		14.6 0.5
. 0	338033		155650	predicted exon	540.6	14
		Y00272	Hs.184572	cell division cycle 2, G1 to S and G2 to	-494.8	1
		BE568452	Hs.5101	protein regulator of cytokinesis 1	-600	i
	338158		. ~~ 101	predicted exon	311.2	31.1
75	327036			predicted exon	351.8	35.2
				•		

	200000	4 1007000	11 440074	FOT		
		AJ227892	Hs.1462/4		180.2	18
	327568			predicted exon	229	22.9
	324801	AW770553	Hs.14553	steroi O-acyttransferase (acyt-Coenzyme	161.2	16.1
	317850	AI681545	Hs 152982	hypothetical protein FLJ13117	-690	1
5		AW043782			126.4	4.5
		A1685464			170.2	17
		X73608		sparc/osteonectin, cwcv and kazal-like d	-80	0
		Al476732			466.8	46.7
	315240	R38772	Hs.172619	KIAA1106 protein	277	27.7
10	338388			predicted exon	267.6	26.8
	338442			predicted exon	256	25.6
	_	AW247252	Hs 75514	nucleoside phosphorylase	1247.8	24.2
	338645	ATTE TI EUL	183.70011	predicted exon	206	20.6
		4150007	11- 400430	• • • • • • • • •		
1.5		N58907	Hs.162430		204.8	20.5
15		BE169746	Hs.12504	hypothetical protein DKFZp761D081	203.6	20.4
	330305			predicted exon	199.8	20
	308248	Al560919		gb:tq41g10.x1 NCI_CGAP_Ut1 Homo sapiens	3	199.4 19.9
		A1833240		gb:at76d10.x1 Barstead colon HPLRB7 Homo		19.8
			He 258188	Homo sapiens cDNA FLJ11674 fis, clone HE		19.1
20						
20		R43240		tumor differentially expressed 1	189.2	18.9
		T91980	Hs.221074		187.6	18.8
	300378	Z45270		hypothetical protein FLJ22672	271.6	18.7
	317478	Al343569	Hs.107000	Homo sapiens mRNA for WDC146, complete	C	187 18.7
	317559	AW452344	Hs.129977	ESTs	184.2	18.4
25		AI873346			182.8	18.3
	334834	10010010	110.211000	predicted exon	178.8	17.9
		Designs				
		D62892		gb:HUM337C07B Clontech human aorta poly/		177.2 17.7
		AL121460	Hs.2/26/3	hypothetical protein FLJ20508	316.4	17.6
	328548			predicted exon	174.6	17.5
30	317108	AA884000	Hs.8173	hypothetical protein FLJ10803	172.4	17.2
	318013	Al188183	Hs.144078	ESTs	326	17.2
		AW382682			170.8	17.1
		AW173339			169.8	17
					169.4	
25		AW975920				16.9
35		AA377578	HS.65234	hypothetical protein FLJ20596	169.2	16.9
	325843			predicted exon	321.4	16.9
	316012	AA764950	Hs.119898	ESTs	1047.2	16.9
	309687	AW236154	Hs.77385	myosin,fightpolypeptide6,alkali,smoothmu	168.2	16.8
	323329	AL134744	Hs.10852		168	16.8
40		W05086	Hs.114256		167.4	16.7
-10		Al422023	Hs.161338		298.6	16.6
		AW977642	HS.291742		165.6	16.6
	338728			predicted exon	165.4	16.5
	316609	AW292520	Hs.122082		165	16.5
45	305989	AA888220		gb:oj15h01.s1 NCI_CGAP_Kid5 Homo saplen	S	164.6 16.5
	312642	AW052128		gb:wx26c02.x1 NCI_CGAP_Kid11 Homo saple	en	164 16.4
	339236			predicted exon	163.6	16.4
		Al217713	He 147586		161.8	16.2
50		AW207582			582.2	16.2
50			Hs.147482		161.2	16.1
		H51696	Hs.89278	hypothetical protein FLJ11186	161	16.1
	317336	AW014637			160	16
	309871	AW300366		gb:xs63b05.x1 NCI_CGAP_Kid11 Homo sapia	า	159.8 16
			Hs.129709	Homo sapiens chromosome 19, cosmid R302	1	159 15.9
55		N52883	Hs.102676		159	15.9
		AW015994		gb:Ul-H-Bl0p-abh-g-09-0-Ul.s1 NCI_CGAP_S		15.9
		N62401		EST	158.6	15.9
		A1684535	Hs.200811		158.4	15.8
~ 0	336059			predicted exon	157.4	15.7
60	302790	AJ245245		gb:Homo saplens mRNA for immunoglobulin	155.8	15.6
	328418			predicted exon	153.8	15.4
	304229	AK000149	Hs.29493	hypothetical protein FLJ20142	153.6	15.4
	331606	AW273285		ESTs	153	15.3
	338962			predicted exon	664.4	15.3
65		Al204202			152.6	15.3
55		MAUNEUL				
	336228	4141000040		predicted exon		15.2
		AW072916		zinc finger protein 131 (done pHZ-10)	152.2	15.2
		AI806867			152.2	15.2
	311943	Al469911	Hs.26498	hypothetical protein FLJ21657	152	15.2
70	314680	Al247425	Hs.152182	ESTs	151.4	15.1
		N29696		EST	151.2	15.1
	338116			predicted exon	151.2	15.1
	329863			predicted exon	150.6	15.1
		AW452886	Ue 230407		149.6	15.7
75						
13		AA868583				15
	331138	R63816	Hs.28445	ESIS	149.6	15

	316561	Al917222	Hs.121655	ESTs	149.4	14.9
	328695			predicted exon	149.2	14.9
	302282	BF396283	Hs.173987	eukaryotic translation initiation factor	148.4	14.8
		F11802	Hs.6818	ESTs	148.2	14.8
5				Homo saplens cDNA FLJ14190 fis, clone NT		14.8
•			Hs.248865			
					147.8	14.8
			Hs.124367		147.8	14.8
			0Hs.82101	,	229	14.8
	324188	AW274439	Hs.252709	ESTs	147.6	14.8
10	317441	AA922798	Hs.196583	ESTs	147.4	14.7
	317584	AI825890	Hs.220513	FSTs	146.8	14.7
		Al308206			146.8	14.7
		AA206045	113.101333		146.6	
				gb:zq77f05.s1 Stratagene hNT neuron (937		14.7
15	313952	F20956		gb:HSPD05390 HM3 Homo saplens cDNA de		146.6 14.7
15			Hs.15713		263.8	14.7
	309195	Al904895	Hs.9614	nucleophosmin (nucleolar phosphoprotein	146.2	14.6
	321860	N47474	Hs.212631	ESTs	146.2	14.6
	330187			predicted exon	146	14.6
		AA463571	Hs.172550	polypyrimidine tract binding protein (he	145.6	14.6
20			Hs.201366		145.2	14.5
20						
				UDP-N-acetylgtucosamine:a-1,3-D-mannosid		14.5
			Hs.133403		144.8	14.5
	302749	M16951		gb:Human Ig mu-chain mRNA VDJ4-region, 5	144.6	14.5
	322357	A1734258	Hs.245367	ESTs, Weakly similar to ALU1_HUMAN ALU :	S	144.6 14.5
25	300391	Al927371	Hs.288839	hypothetical protein FLJ12178	144.4	14.4
	326077			predicted exon	144.4	14.4
		Y18264	He 122004	sal (Drosophila)-like 1	144	14.4
			Un 440047	Versions of the Court for since t		
				Homo sapiens cDNA: FLJ23077 fis, clone L	144	14.4
20		T88693	Hs.226410		144	14.4
30		Al969727		ESTs	143.2	14.3
	305159	AA659166	Hs.275668	EST, Weaklysimilarto EF1D_HUMANELONGA	TIONF	143 14,3
				ribosomalproteinL13a	142.8	14.3
			Hs.303527		142.8	14.3
			Hs.122139		142.8	14.3
35				basic-helix-loop-helix-PAS protein		
"					142.6	14.3
		A1824879		ESTs, Weakly similar to 1207289A reverse	142.2	14.2
			Hs.293565	ESTs, Weakly similar to putative p150 [H	142.2	14.2
	332283	R40855	Hs.100839	EST	142	14.2
	302894	AA719572	Hs.274441	Homo sapiens mRNA; cDNA DKFZp434N011	(fr	141.2 14.1
40		R35391	Hs.252831		141	14.1
		Al204491	Hs.151502		141	14.1
			Hs.122872			
					140.8	14.1
			Hs.150276		140.8	14.1
45		N45600	Hs.46534	Homo sapiens mRNA; cDNA DKFZp434P0714		140.8 14.1
45	327365			predicted exon	140.8	14.1
	331264	AA278898	Hs.225979	hypothetical protein similar to small G	140.8	14.1
	324545	AW501944	Hs.127243	Homo sapiens mRNA for KIAA1724 protein,	140.4	14
		AA211586		gb:zn56d05.s1 Stratagene muscle 937209 H		14
			Hs.149065		140.2	14
50		BE247449				14
50			115.5 1002	hypothetical protein FLJ10525	140.2	
		AA595765		gb:nj28g06.s1 NCI_CGAP_AA1 Homo sapien:		139.814
			Hs.202450		139.8	14
			Hs.146840		139.8	14
	324137	AA393127	Hs.222762	ESTs	139.8	14
55	328262			predicted exon	139.6	14
	322349	AK001279	Hs.180171	Homo sapiens cDNA FLJ10417 fis, clone NT		13.9
			Hs.130865		139.4	13.9
		AA059387		gb:zf66d01.s1 Soares retina N2b4HR Homo		13.9
			Un DOFESE	hypothetical protein PRO2955		
60		A1143 1433			139.2	13.9
UU	335946			predicted exon	139.2	13.9
			Hs.132133		138.8	13.9
	313796	Al797169	Hs.208486	ESTs	138.6	13.9
	333977			predicted exon	138.6	13.9
	324845	AW969635	Hs.283718		138.2	13.8
65	331139			gb:yi16g12.s1 Soares placenta Nb2HP Homo		13.8
	331131				669.6	13.8
	321250		Hs.151692		138	13.8
				ESTs, Weakly similar to ALU1_HUMAN ALU S		137.8 13.8
=-	331252	W52470	Hs.34578	alpha2,3-sialyltransferase	137.8	13.8
70	337407			predicted exon	137.8	13.8
		AW512014		gbxx68a03.x1 NCL_CGAP_Lym12 Homo sapi		137.4 13.7
			Hs.188817		137.4	13.7
	327373				137.2	13.7
		A A 224 E 04	Hs.304123	<u></u>		
75					136.6	13.7
13			Hs.120260		136.4	13.6
	315231	AA705809	Hs.119922	E918	136.2	13.6

	319500	T39310	Hs.1139	éald chaek domain nortain A	420.0	42 C
	–			cold shock domain protein A	136.2	13.6
		AW969706	rts.290302		136.2	13.6
	328937			predicted exon	136.2	13.6
_	329073			predicted exon	136.2	13.6
5		AV659082			136	13.6
	311992	AL360200	Hs.114145	ESTs	135.8	13.6
	316497	AA766457	Hs.136849	ESTs	135.8	13.6
	317677	AA968594	Hs.127868	ESTs	135.8	13.6
	321680	W02848	Hs.93704	ESTs	135.8	13.6
10	326080			predicted exon	135.8	13.6
		AF036943	Hs.172619	KIAA1106 protein	135.8	13.6
				ribosomal protein, large P2	135.6	13.6
		Al223207			135.6	13.6
		AW449382			135.6	
15						13.6
13		R15138	MS. 1000/U	Homo sapiens clone 25052 mRNA sequence		13.5
		AA884104			134.8	13.5
		N58198			134.8	13.5
		H09604	Hs.13268		134.6	13.5
	332776	AF241850	Hs.151428	ret finger protein 2	134.4	13.4
20	332494	AA282330	Hs.145668	ESTs	134.2	13.4
	334376			predicted exon	134.2	13.4
	313264	N93416	Hs.118228		133.6	13.4
		AA351109		Tax1 (human T-cell leukemia virus type I	133.2	13.3
		T87398	Hs.205816		132.6	13.3
25	310354	AA993807	He 167367	ECTe	132.6	13.3
23		Al242106	1 13. 107 307	gb:qh92a02.x1 Soares_NFL_T_GBC_S1 Hom		132.2 13.2
			U- 204402			
				Apg12 (autophagy 12, S. cerevisiae)-like	131.8	13.2
		AI274963			131.2	13.1
20		AW510641			220.6	13
30		F12650	Hs.13287	ESTs	125.4	12.5
	329451			predicted exon	123.4	12.3
	337603			predicted exon	572	12.2
	312480	R68651	Hs.144997	ESTs	121.4	12.1
	324934	AW452051	Hs.147546	ESTs	119.4	11.9
35		BE178025		hypothetical protein FLJ20080	117	11.7
-		Al792566	1,0.1012	gb:qi74f02.y5 NCI_CGAP_Ov26 Homo saplen		116.6 11.7
			He 283055	Homo sapiens clone GLSH-2 similar to gli	112.8	11.3
		BE003191			112.6	11.3
		AA614406	ns.119000			112.4 11.2
40			11- 007707	gb:np46f05.s1 NCI_CGAP_Br11 Homo sapier		
40		Al139253		zinc finger protein 41	111.2	11.1
		AA347945			111	11.1
				ribosomalproteinS4,X-linked	111	11.1
		AW419225			110.2	11 -
4 10		AW452334			110.2	11
45		W49701	Hs.29667		109.4	10.9
	315099	AA806536	Hs.291841	ESTs	109	10.9
	304793	AA583264	Hs.182979	ESTs ribosomalproteini.12 k(AA1238 portein	108.8	10.9
	330815	AA019211	Hs.236463	KIAA1238 protein	108.8	10.9
	304044	T81656	Hs.252259		714.8	10.8
50	325222			predicted exon	135	10.8
• •	325889				814.6	10.8
		AW891130	He 38173	ESTs	107.8	10.8
		AA496212			106.2	10.6
	302330	A1476803	113.100102	gb:tj77e12.x1 Soares_NSF_F8_9W_OT_PA_I		270.6 10.6
55		Al301041	Un 450474	go.greez.x coales_rol _ o_ori_or_ro_	106	10.6
55						
		AW157431			233	10.6
	318787		Hs.22657	ESTs	105.8	10.6
		Al927226	Hs.175610		105.2	10.5
	326788			predicted exon	104.4	10.4
60		AA830640			104	10.4
	314482	AW085525	Hs.134182	ESTs	234	10.4
	323597	Al185693	Hs.135119	ESTs	102.4	10.2
	302623	AW836724	Hs.194110	hypothetical protein PRO2730	162.4	10.2
		AI791531			101	10.1
65	324315			zinc finger protein 265	100.2	10
		AA256465			99.2	9.9
		AA554913			98.2	9.8
	327876	70.001010	I IS. TULEST	predicted exon	98.2	9.8
		D47424	Un CCEA		98	
70		R17424	Hs.6650	vacuolar protein sorting 45B (yeast homo		9.8
70	327747			predicted exon	97.6	9.8
	327844				97.4	9.7
		Al061192	Hs.166517		97.2	9.7
	329414				97.2	9.7
~~		A1089667	Hs.270713			9.7
75	307010	Al140014		gb:qa68f09.x1 Soares_fetal_heart_NbHH19W		9.7
	319792	Al138635	Hs.22968		385.4	9.6

		AA811688	Hs.82113		96	9.6
	329440			predicted exon	93.8	9.4
	310381	Al263059	Hs.145594	ESTs	93.4	9.3
	318824	F05771	Hs.27226	ESTs	93.4	9.3
5	328957			predicted exon	92.2	9.2
	318804	Z42549	Hs.160893		92	9.2
	330836	AA055611		ESTs, Moderately similar to ALU4_HUMAN A	92	9.2
			Hs.325708		91.8	9.2
			Hs.254333		91.4	9.1
10			16.234333			
10		H86161		gb:ys94b01.r1 Soares retina N2b5HR Homo		9.1
	330306			predicted exon	91	9.1
				regulator of G-protein signalling 17	90	9
	313275	A1027604	Hs.159650	ESTs	110.4	8.8
	302593	H54855	Hs.36958	ESTs ·	88	8.8
15	321421	BE465115	Hs.171688	ESTs	86.2	8.6
	330832	AJ133530	Hs.62930	ESTs	456.4	8.6
			Hs.297260		86	8.6
				Homo sapiens cDNA FLJ14080 fis, clone HE		8.6
	328688	500000	. 15.50 1555	predicted exon	85.6	
20	325251			predicted exon	85.4	8.5
20						
	329088	11770007		predicted exon	85.4	8.5
		W79027	Hs.271762		84	8.4
	337953			predicted exon	451	8.3
	323529	AA284397	Hs.201485	Homo sapiens done FLC0664 PRO2866 mRI		82.6 8.3
25	307041	Al144243		gb:qb85b12.x1 Soares_fetal_heart_NbHH19V	٧	306.88.2
	318285	Al332454	Hs.158412		81.4	8.1
			Hs.14041		81	8.1
	329350			predicted exon	81	8.1
	326169			predicted exon	80.4	8
30	338038			predicted exon	1024.2	7.9
50		41044540	U- 440204	•		
		Al214510	Hs.146304		77.4	7.7
		D60076		gb:HUM084E10A Clontech human fetal brain		7.7
				solute carrier family 7, (cationic amino	76	7.6
		Al470235	Hs.172698	= - :	150.6	7.5
35	315650	AA649042	Hs.269615	ESTs	73.4	7.3
	324328	AA447276	Hs.292020	ESTs	210.4	7.1
	332622	R10674	Hs.128856	CSR1 protein	70.2	7
	328229			predicted exon	69.4	6.9
		175260	Hs.98321	hypothetical protein FLJ14103	68.6	6.9
40		Al187742	Hs.125562		308.6	6.9
70		AW515800	115.123302			
			Un 440500	gb:hd88g01.x1 NCI_CGAP_GC6 Homo sapie		67.8 6.8
		AAU17000	HS. 14000U	enolase 2, (gamma, neuronal)	377.2	6.7
	325753			predicted exon	105.2	6.6
4.5		Al004377	Hs.200360	Homo sapiens cDNA FLJ13027 fis, clone NT		6.4
45	302592	AA294921	Hs.250811	v-ral simian leukemia viral oncogene hom	361.6	6.3
	314578	AA410183	Hs.137475	ESTs	201.6	6.1
	335986			predicted exon	108.6	6
	321478	AW402593	Hs.123253	hypothetical protein FLJ22009	528 ·	6
		AA666019		gb:ag44a04.s1 Jia bone marrow stroma Hom		5.9
50		AA070605		gb:zm53h09.s1 Stratagene fibroblast (937	78.6	5.6
-		AJ235667		gb:Homo sapiens mRNA for immunoglobulin		5.5
			U. 77050			
	301976		Hs.77256	enhancer of zeste (Drosophila) homolog 2	479.2	5.4
	310021	AW293399	ns. 144904	nuclear receptor co-repressor 1	792.4	5.3
<i></i>				BENE protein	2423.8	5.3
55			Hs.176101		523.2	5.1
			Hs.172445		578	5.1
	303190	BE280787	Hs.16079	hypothetical protein FLJ10233	223	5.1
	315587	AI268399	Hs.140489	ESTs	136.2	5
	333122			predicted exon	399	5
60		Al220072	Hs.165893		234.4	4.9
				transcription factor EC	68	4.9
				EST, Weakly similar to A27217 glucose tr	258.8	4.8
		Z44067	ns. 1033/	ESTS	376.6	4.8
65	327220			predicted exon	47.4	4.7
65			Hs.179872		177.4	4.7
			Hs.151072		205.2	4.6
	313339			Homo sapiens dDNA FLJ13608 fis, done PL		4.5
	318634	T49598	Hs.156832	ESTs	475.2	4.5
			Hs.278679	a disintegrin and metalloproteinase doma	388.6	4.4
70					.81.6	4.4
		AW051913		gb.wx24a09.x1 NCI_CGAP_Kid11 Homo sapi		102.4 4.3
		AA889992	He 21RE	eukaryolictranslationelongationfactor1ga	451.2	4.2
		~~~03334	i io.4 100			
	330020	VITE CATTY	U= 04070	predicted exon	61.2	4.1
75			ms.913/9	ribosomal protein L26	342	3.9
75		AW979268		gb:EST391378 MAGE resequences, MAGP H		56.4 3.8
	J15131	AI/53709	Hs.152484	E918	130.4	3.7

						••	
		Al493591	Hs.78146	platelet/endothelial cell adhesion molec	3179.6	3.6	
	333585	H93366	Hs.7567	predicted exon Homo sapiens cDNA: FLJ21962 fis, clone H	175.4	3.5 3.5	
				Homo sapiens cell recognition molecule C	350.2	3.4	
5		R71072			322.8	3	
•		Z44203	Hs.26418		25	2.5	
		Al123705			773.4	2.5	
	324181	AI025476	Hs.131628	ESTs	634.8	2.4	
		AW205369			54.2	2.4	
10				transcription factor BMAL2	23.4	2.3	
	308852	AI829848	Hs.182937	peptidytprolytisomeraseA(cyclophilinA)	92	2.3	
				Homo sapiens cDNA FLJ13585 fis, clone PL		2.3	
				DKFZP434N178 protein	76.2	2.2	
15		N24236		nucleosome assembly protein 1-like 1	253.2 136.6	2.1 2.1	
13				homeo box B4 NADPH oxidase 4	720	1.8	
				ESTs, Weakly similar to 1605244A erythro	27	1.8	
		AW196663			303.8	1.6	
		W35132	Hs.267442		189	1.5	
20				translocase of inner mitochondrial membr	14.4	1.4	
				Mitochondrial Acyl-CoA Thioesterase	529.8	1.4	
	333123			predicted exon	396.2	1.4	
	328455			predicted exon	91.8	1.3	
	334458			predicted exon	406.4	1.3	
25		AA643008			413.4	1.1	
		AW338564			-30,8	1	
				Homo saplens cDNA FLJ12981 fis, clone NT		1	
				coagulation factor II (thrombin) recepto	-73.6	1	
20		AW367295			-43.8	1	
30				ESTs, Weakly similar to ALU4_HUMAN ALU \$		-63 1	
		AW275110			-67 -395.2	1	
		AI792140 AL121278			-1.6	1	
		AW975183			4.4	i	
35		AW970939			-282.8	1	
-	333610			predicted exon	-152.6	1	
	335093			predicted exon	-23.2	1	
	339403			predicted exon	-331.2	1	
	302820	X04588	Hs.85844		591.2	1	
40		R56151	Hs.93589			276.6 1	
		AW300094	Hs.136252	and the second s	135	0.9	
	326946	D=444000	11 420004	predicted exon	727.4	0.9	
				ESTs, Weakly similar to P4HA_HUMAN PROL		122.8 0.9	
45		AKUU12/U	HS. 196086	hypothetical protein FLJ10408	304	0.9	
43	329732	DE624907	Un 2227	predicted exon transmembrane 4 superfamily member 1	109.2 414.8	0.9 0.7	
	333121	BE621807	U2:2221	predicted exon	87.8	0.7	
	333120			predicted exon	379.8	0.7	
		AW797956	He 75748	proteasome (prosome, macropain) subunit,	589.2	0.7	
50		AA769365			-87	0.6	
-		BE409857			347.4	0.6	
	333169			predicted exon	-1182	0.6	
	335095			predicted exon	106.4	0.6	
	335815			predicted exon	-156	0.6	
55	330232			predicted exon	102.6	0.6	
				ESTs, Moderately similar to ALU5_HUMAN A		0.5	
		F04225 .			-14.6	0.5	
				NESH protein	267.6	0.5	
60		AA432067			85	0.5	
60				homolog of mouse C2PA	-70 -195.4	0.4 0.4	
		A1077462		ribosomalprotein,large,P0	-33.4	0.4	
		N46436			-392	0.4	
	332961	1410100	113, 10322.1	predicted exon	-5.6	0.4	
65	322796	W31178	Hs.154140	Homo sapiens ovary-specific acidic prote	-880.6	0.3	
•••	328857			predicted exon	55.2	0.3	
		AA743935	Hs.202329		43.4	0.3	
		AM720102	Hs.267596	ESTs	-180.4	0.3	
	331263	MW/ OU 132					
<b>7</b> 0	335987			predicted exon	-134	0.3	
70	335987 311923	T60843	Hs.189679	ESTs	12.2	0.3	
70	335987 311923 310522	T60843 AW134529	Hs.189679 Hs.244647	ESTs ESTs	12.2 -187.8	0.3 0.3	
70	335987 311923 310522 315363	T60843 AW134529 AA759190	Hs.189679 Hs.244647 Hs.121454	ESTs ESTs ESTs, Weakly similar to olfactory recept	12.2 -187.8 80	0.3 0.3 0.3	
70	335987 311923 310522 315363 302032	T60843 AW134529 AA759190 NM_001992	Hs.189679 Hs.244647 Hs.121454 2Hs.128087	ESTs ESTs, Wealdy similar to olfactory recept coagulation factor II (thrombin) recepto	12.2 -187.8 80 -877	0.3 0.3 0.3 0.3	
	335987 311923 310522 315363 302032 313140	T60843 AW134529 AA759190 NM_001992 BE265133	Hs.189679 Hs.244647 Hs.121454 2Hs.128087 Hs.217493	ESTs ESTs, Wealdy similar to olfactory recept coagulation factor II (thrombin) recepto annexin A2	12.2 -187.8 80 -877 95.4	0.3 0.3 0.3 0.3 0.3	
70 75	335987 311923 310522 315363 302032 313140 310860	T60843 AW134529 AA759190 NM_001992 BE265133 AW015920	Hs.189679 Hs.244647 Hs.121454 2Hs.128087 Hs.217493 Hs.161359	ESTs ESTs, Wealdy similar to olfactory recept coagulation factor II (thrombin) recepto annexin A2	12.2 -187.8 80 -877 95.4 -239	0.3 0.3 0.3 0.3	0.3

	328520			predicted exon	-109.2	0.2
		NISA 04200	0U- 2110EE	CD3-epsilon-associated protein; antisens		0.2
	244904	AIRCCOM	113.Z I 1500	Upper content of NA CI 140440 for store NA	10	
	311004	A1600921	NS.203349	Homo saplens cDNA FLJ12149 fis, clone MA		0.2
5	315065	AKUU1122	HS.105859	hypothetical protein FLJ10260	-46.2	0.2
5		AA228366	Hs.115122		-308.8	0.2
	335697			predicted exon	-47.2	0.2
	335989			predicted exon	89	0.2
	320606	AW867943	Hs.127216	hypothetical protein FLJ13465	-205.6	0.2
	329745			predicted exon	103	0.2
10	313628	AW419069	Hs.209670	ESTs	-177.8	0.2
	334616			predicted exon	-936.6	0.2
		AI821267	Hs.207243		-7.2	0.2
		AI026984	Hs.293662		-18.4	0.2
	335211	A1020304	113.230002	predicted exon	-142	0.2
15		AA375957	Un CCOD	ESTs	-100	0.1
13			N\$.000Z	=		
		AW452904	11- 404700	gbtUH+Bi3-aly-h-11-0-Ul.s1 NCI_CGAP_Su		0.1
		AIU15862	Hs.131793		-250.6	0.1
	332833			predicted exon	-374.2	0.1
20			Hs.36566		-27.6	0.1
20			Hs.187820		-288.2	0.1
			Hs.208558		-735.6	0.1
	330775	AW247020	Hs.250747	SUMO-1 activating enzyme subunit 1	53.6	0.1
	333168			predicted exon	-1041.8	0.1
	332079	Al308876	Hs.103849	ESTs	19.4	0.1
25	322724	AF161442	Hs.191591	Homo sapiens HSPC324 mRNA, partial cds	-123.6	0.1
			Hs.64341		-46.4	0.1
				DC2 protein	-156.4	0.1
			Hs.171532		-146.6	0.1
			Hs.163780		-304	0.1
30			Hs.183817		-445	0.1
-		R43707		ESTs, Weakly similar to PIHUSD salivary	-111.8	0.1
		Al832519	1 55.100 100	gb:at69h03.x1 Barstead colon HPLRB7 Homo		0
			Hs.207144		-62.6	ŏ
				2,3-bisphosphoglycerate mutase	-15	Ď
35			Hs.42699		-1059.6	ŏ
22					-332.2	0
			MS. 102090	MCT-1 protein		
		AA961367	11- 74000	gb:or52a05.s1 NCI_CGAP_GC3 Homo sapier		21.8 0
		Al292148		SWI/SNF related, matrix associated, acti	-97.2	0
40					-2222	0
40			Hs.252844		-881.6	0
		AW288085		hypothetical protein	28.4	0
		AF167706	Hs.19280	-,	-722	0
	335990			predicted exon	-421	0
		AB033888			35.4	0
45	316820	Al627912	Hs.130783	Forssman synthetase	-373.6	0
	337429			predicted exon	-257	0
	331192	BE622021	Hs.152571	ESTs, Highly similar to IGF-II mRNA-bind	-33	0
	330609	Al346201	Hs.76118	ubiquitin carboxyl-terminal esterase L1	-280	0
	323593	AI739435	Hs.39168	ESTs	-3627.6	0
50	302704	AA531133	Hs.4253	hypothetical protein MGC2574	-278.6	0
		NM_004579		71	-244	Ō
	332374				-1204.2	Ō
	333221				-189.6	ŏ
	335988			F	-122.6	0
55	•••	AI984144	Hs.66713		-2257.4	0
J.J		BE621697			-359.2	0
		AF131781			-874.6	0
	337113	VL 121101	113,04733		-074.0 -24.6	0
	335149				-24.6 -191.8	0
	JJU 143			hierioren evrit	-131.0	U

#### TABLE 6A

Table 6A shows the accession numbers for those pkeys lacking unigeneiD's for Table 6. The pkeys in Table 7 lacking unigeneiD's are represented within Tables 1-6A. For each probeset we have listed the gene cluster number from which the oligonucleotides were designed. Gene clusters were compiled using sequences derived from Genbank ESTs and mRNAs. These sequences were clustered based on sequence similarity using Clustering and Alignment Tools (DoubleTwist, Oakland California). The Genbank accession numbers for sequences comprising each cluster are listed in the "Accession" column.

10

Unique Eos probeset identifier number

Pkey: CAT number:

Gene cluster number Genbank accession numbers

Accession:

328229 c_6_hs

328262 c_6_hs

75

```
Pkey CAT Number Accession
         320925 1525201_1
                               D62892 D79755 D62760
20
         321614 87866_1
                                H86161 AA054308 AA018955
         313952 136885_1
                               F20956 AA129374 AA133740 AW819878
                                AW979268 AA878419 AA431342 AA431628
         314648 293660_1
                                M16951 M16952 M16948 M16949 M16950
         302749 458_107
         312362 764066_1
                               AW015994 R39898 AW000978 Al598202 Al521706
25
         312542 1522649_1
                               D60076 D60259 D61037
         312642 1005225 1
                               AW052128 H51439 H51481
         312986 171879_1
                               AA211586 F35799 AA211641 F29720 AW937387 AW937408
         329350 c_x_hs
         329414 c_y_hs
30
         329440 c_y_hs
        329451 c.y.hs
338033 CH22_6528FG_LINK_EM:AC00
338038 CH22_6535FG_LINK_EM:AC00
338116 CH22_6650FG_LINK_EM:AC00
35
         338158 CH22_6700FG__LINK_EM:AC00
         329732 c14_p2
         329745 c14_p2
308106 Al476803
         329863 c14_p2
40
         338316 CH22_6944FG_LINK_EM:AC00
         308248 Al560919
         338388 CH22_7034FG_LINK_EM:AC00
        338442 CH22_7109FG_LINK_EM:AC00
338645 CH22_7410FG_LINK_EM:AC00
338728 CH22_7527FG_LINK_EM:AC00
45
         308877 Al832519
         338962 CH22_7838FG__LINK_DJ32I10
         308886 AI833240
        333120 CH22_349FG_81_3_LINK_EM:A
333121 CH22_350FG_81_4_LINK_EM:A
333122 CH22_351FG_81_6_LINK_EM:A
50
         333123 CH22_352FG_81_7_LINK_EM:A
         333168 CH22_400FG_94_1_LINK_EM:A
         333169 CH22_401FG_94_2_LINK_EM:A
55
         333221 CH22_458FG_105_1_LINK_EM:
         326077 c17_hs
         326080 c17_hs
         326169 c17_hs
         326198 c17_hs
60
         326230 c17_hs
         333585 CH22_846FG_203_4_LINK_EM:
         333610 CH22_871FG_217_5_LINK_EM:
335093 CH22_2423FG_492_3_LINK_EM
         335095 CH22_2425FG_492_5_LINK_EM
65
         335149 CH22_2484FG_499_5_LINK_EM
         326759 c20_hs
333977 CH22_1254FG_309_6_LINK_EM
         326788 c20_hs
         335211 CH22_2550FG_511_2_LINK_EM
70
         305192 AA666019
         303973 AW512014
         303992 AW515800
         326946 c21_hs
```

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328418 c_7_hs
         328455 c_7_hs
         335697 CH22_3058FG_596_12_LINK_E
         328520 c_7_hs
         328548 c_7_hs
         335815 CH22_3187FG_618_3_LINK_EM
         328688 c_7_hs
         328695 c_7_hs
         307010 Al140014
         337113 CH22_5058FG_493_1_
10
         307041 Al144243
         328700 c_7_hs
         335946 CH22_3324FG_646_20_LINK_D
         33598 CH22_3366FG_654_10_LINK_D
335987 CH22_3367FG_654_11_LINK_D
335988 CH22_3368FG_654_12_LINK_D
335989 CH22_3369FG_655_2_LINK_DJ
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         335990 CH22_3370FG_655_4_LINK_DJ
337214 CH22_5288FG_613_7_
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         330020 c16_p2
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         328857 c_7_hs
         328937 c_8_hs
         328957 c_8_hs
330187 c_4_p2
25
         337407 CH22_5607FG_755_1_
337429 CH22_5633FG_762_3_
         330232 c_5_p2
         307414 Al242106
30
         330305 c_7_p2
         330306 c_7_p2
337603 CH22_5896FG__LINK_C20H12.
         337953 CH22_6395FG_LINK_EM:AC00
339236 CH22_8181FG_LINK_BA354I1
35
         339403 CH22_8384FG__LINK_BA232E1
         309349 AW051913
         325222 c10 hs
         325251 c10_hs
                                AI792566 AI053836 AI054127 AI792489 AI288324
         318188 956161_1
40
         309871 AW300366
         325544 c12_hs
         309931 AW341683
         332833 CH22_50FG_17_7_LINK_C20H1
302779 33837_1 AJ235667 AJ23566
                                AJ235667 AJ235666 AJ235664 AJ235665 AJ235668 AJ235669 AJ235670
                                AJ245245 AJ245247 AJ245257 AJ245248 AJ245254 AJ245256 AJ245253 AJ245203 AJ245250 AJ245252 AJ245243 AJ245204
45
         302790 34168_1
                                AJ245201 AJ245206 AJ245246 AJ245255 AJ245205 AJ245202 AJ245251 AJ245249 AJ245207 AJ245244
         332961 CH22_185FG_48_18_LINK_EM:
         325753 c14_hs
         327036 c21_hs
         325843 c16_hs
50
         325889 c16_hs
         304261 AA059387
         304275 AA070605
         334376 CH22_1670FG_379_8_LINK_EM
         327220 c_1_hs
304363 AA206045
55
         334458 CH22_1757FG_391_2_LINK_EM
         327365 c_1_hs
         327373 c_2_hs
60
         334616 CH22_1923FG_411_15_LINK_E
         327414 c_2_hs
         327568 c.3_hs
336034 CH22_3419FG_678_5_LINK_DJ
336059 CH22_3445FG_684_2_LINK_DJ
65
         334834 CH22_2148FG_439_3_LINK_EM
         304782 AA582081
         304876 AA595765
         327747 c_5_hs
         336228 CH22_3626FG_730_4_LINK_DA
70
         329073 c.x.hs
         329088 c_x_hs
         304969 AA614406
         327844 c_5_hs
         327876 c_6_hs
75
         306352 AA961367
         331131 genbank_R54797
                                           R54797
```

331139 genbank_R65706 R65706 331420 675963_1 AW452904 AW449414 BE467906 Al298565 BE549932 BE326357 F04362

#### **TABLE 6B**

Table 6B shows the genomic positioning for those pkeys lacking unigene ID's and accession numbers in Table 6. The pkeys in Table 7 lacking unigeneID's are represented within Tables 1-6B. For each predicted exon, we have listed the genomic sequence source used for prediction. Nucleotide locations of each predicted exon are also listed. 5

Pkey: Ref: Unique number corresponding to an Eos probeset
Sequence source. The 7 digit numbers in this column are Genbank Identifier (GI) numbers. "Dunham I. et al." refers to the publication entitled "The DNA" 10 sequence of human chromosome 22.* Dunham I. et al., Nature (1999) 402:489-495. Indicates DNA strand from which exons were predicted. Strand:

Nt_position: Indicates nucleotide positions of predicted exons.

15				
	Pkey	Ref	Strand	Nt_position
	332961	Dunham, I. et.al.	Plus	2521424-2521555
20	333221	Dunham, l. et.al.	Plus	3978070-3978187
		Dunham, I. et.al.	Plus	6234778-6234894
	333610		Płus	6547007-6547116
	334376		Plus	13902218-13902331
	334458	Dunham, I. et.al.	Phis	14353496-14353572
25	334616	Dunham, I. et.al.	Plus	15176123-15176470
		Dunham, I. et.al.	Plus	21497441-21497587
	335211	Dunham, I. et.al.	Plus	21774611-21774680
	335697	Dunham, I. et.al.	Plus	25481456-25481649
20		Dunham, I. et.al.	Plus	27967791-27967852
30	335987	Dunham, I. et al.	Plus	27971413-27971481
		Dunham, I. et.al.	Plus	27977912-27978013
		Dunham, I. et.al.	Plus	27983788-27983860
		Dunham, I. et al.	Plus	27988532-27988608
25	336034	Dunham, I. et.al.	Plus	29014404-29014590
35	337953		Plus	6827029-6827125
	338033		Plus	8092128-8092271
	338038		Plus	8138219-8138392
		Dunham, I. et al.	Plus	17089711-17089988
40		Dunham, I. et.al.	Pius	19980640-19980698 29581892-29582020
40	332833	Dunham, I. et al.	Plus	1119848-1119705
	333120	Dunham, I. et.al. Dunham, I. et.al.	Minus Minus	3307508-3307427
	333121	Dunham, I. et al.	Minus	3308446-3308358
	333122	Dunham, I. et.al.	Minus	3309596-3309531
45	333123		Minus	3310817-3310749
	333168	Dunham, I. et al.	Minus	3729896-3729788
	333169		Minus	3730864-3730767
	333977	Dunham, I. et al.	Minus	8722928-8722725
	334834	Dunham, I. et al.	Minus	17182681-17182535
50	335093		Minus	21297367-21297214
	335095	Dunham, I. et.al.	Minus	21292546-21292381
	335815	Dunham, I. et.al.	Minus	26320518-26320421
	335946	Dunham, I. et.al.	Minus	27487203-27487035
	336059	Dunham, I. et.al.	Minus	29184079-29183969
55	336228	Dunham, I. et.al.	Minus	30904602-30904497
	337113	Dunham, i. et.al.	Minus	21233344-21233237
	337214	Dunham, I. et.al.	Minus	26095902-26095502
	337407	Dunham, I. et.al.	Minus	31886652-31886567
60	337429	Dunham, I. et.al.	Minus	32086238-32086079
60	337603	Dunham, I. et al.	Minus	1299296-1299194
	338116	Dunham, I. et.al.	Minus	10614071-10613814
	338158	Dunham, I. et.al.	Minus	11794465-11794343
	338388 338645	Dunham, I. et.al.	Minus	18662403-18662305 24063839-24063775
65	338728	Dunham, I. et.al. Dunham, I. et.al.	Minus Minus	25949039-25948927
05	339236	Dunham, I. et al.	Minus	32773355-32773202
	339403	Dunham, I. et al.	Minus	34050728-34050625
	325222	6525287	Minus	22332-22473
	325251	6682448	Minus	411693-411751
70	325544	6682452	Plus	171228-171286
. •	325753	6682474	Plus	398512-398621
	329745	6065779	Phus	174774-175142
	329732	6065783	Phus	161252-161322
	329863	6691797	Plus	196801-196971
75	325889		Plus	223829-223891
	-			

	325843	6552453	Minus	7126-7232
	330020	6671887	Plus	172397-172491
	326198	5867215	Minus	80295-80674
	326230	5867230	Minus	301868-301972
5	326169	5867255	Minus	128321-128388
	326077	6682495	Minus	312108-312168
	326080	6682495	Plus	478644-478847
	326759	6249610	Plus	97216-97311
	326788	6682503	Plus	277132-277335
10	326946	6004446	Minus	116677-116967
	327036	6531965	Plus	319951-320040
	327220	5867525	Minus	65701-65781
	327365	6552412	Minus	118133-118198
	327414	5867750	Plus	102461-102586
15	327373	5867792	Minus	8186-8742
	327568	5867811	Minus	46152-46287
	330187	6706138	Plus	212923-213020
	327747	5867947	Plus	115322-115498
	327844	6249582	Minus	18895-18958
20	330232	6013526	Plus	113655-113830
	328229	5868105	Minus	120936-121053
	327876	5868140	Plus	103882-104034
	328262	6381906	Plus	11867-12027
	328688	5868262	Plus	626030-626094
25	328700	5868264	Plus	764089-764203
	328695	5868264	Plus	318632-318695
	328418	5868409	Minus	258811-258894
	328455	5868431	Ptus	385576-385633
	328520	••••	Plus	1942075-1942246
30	328548	5868487	Plus	72301-72397
	328857	6381927	Minus	80557-81051
	330305	4877982	Minus	52269-52365
	330306	4877982	Plus	96161-96233
	328937	5868500	Minus .	1448241-1448333
35	328957	6456773	Plus	219195-219297
	329073	5868596	Plus	37838-37956
	329088	5868608	Plus	116738-116950
	329350	6456785	Plus	98911-98969
40	329414	5868874	Plus	942555-942643
40	329440	5868885	Plus	21943-22063
	329451	5868887	Plus	25974-26048

# TABLE 7:

Table 7 depicts Seq ID No., UnigeneID, UnigeneTitle, Pkey, and ExAcon for all of the sequences in Table 8. Seq ID No links the nucleic acid and protein sequence information inTable 8 to Table 7. 5

Unique Eos probeset identifier number Exemplar Accession number, Genbank accession number Unigene number Unigene gene title Sequence Identification Number found in Table 8

Pkey: ExAcon: UnigeneID: Unigene Title: Seq.ID.No.: 10

15	PKey	ExAcon	Unigene ID	Unigene Tiltle	SEQ ID NO
	101545	BE246154	Hs.154210	endothelial differentiation, sphingolipi	Seq ID 1 & 2
	115819	AA486620	Hs.41135	endomucin-2	Seq ID 3 & 4
	424503	NM_002205	Hs.149609	integrin, alpha 5 (fibronectin receptor,	Seq ID 5 & 6
	102917	AI016712	Hs.287797	integrin, beta 1 (fibronectin receptor,	Seq ID 7 & 8
20	102915	X07820	Hs.2258	matrix metalloproteinase 10 (stromelysin	Seq ID 9 & 10
	105330	AW338625	Hs.22120	ESTs	Seq ID 11 & 12
	107385	NM_005397	Hs.16426	podocalyxin-like	Seq ID 13 & 14
	102024	AA301867	Hs.76224	EGF-containing fibulin-like extracellula	Seq ID 15 & 16
25	102024	AA301867	Hs.76224	EGF-containing fibulin-like extracellula	Seq ID 17 & 18
25	134416	X68264	Hs.211579	metanoma cell adhesion molecule	Seq ID 19 & 20
	103036	M13509	Hs.83169 Hs.22575	matrix metalloproteinase 1 (interstitial B-cell CLL/lymphoma 6, member B (zinc fi	Seq ID 21 & 22 Seq ID 23 & 24
	104865 106124	T79340 H93366	Hs.7567	Homo saplens dDNA: FLJ21962 fis, clone H	Seq ID 25 & 26
	109001	AI056548	Hs.72116	hypothetical protein FLJ20992 similar to	Seq ID 27 & 28
30	104764	AI039243	Hs.278585	ESTs	Seq ID 29 & 30
	133200	AB037715	Hs.183639	hypothetical protein FLJ10210	Seq ID 31 & 32
	105263	AW388633	Hs.6682	solute carrier family 7, (cationic amino	Seq ID 33 & 34
	102892	BE440042	Hs.83326	matrix metalloproteinase 3 (stromelysin	Seq ID 35 & 36
	109456	AW956580	Hs.42699	ESTs	Seq ID 37 & 38
35	110906	AA035211	Hs.17404	ESTs	Seq ID 39 & 40
	119073	BE245360	Hs.279477	ESTs	Seq ID 41 & 42
	132050	Al267615	Hs.38022	ESTs	Seq ID 43 & 44
	132490	NM_001290	Hs.4980	LIM domain binding 2	Seq ID 45 & 46 Seq ID 47 & 48
40	102283 101714	AW161552 M68874	Hs.83381 Hs.211587	guantne nucleotide binding protein 11 phospholipase A2, group IVA (cytosolic,	Seq ID 49 & 50
+0	133975	C18356	Hs.295944	tissue factor pathway inhibitor 2	Seq ID 51 & 52
	106793	H94997	Hs.16450	ESTs	Seq ID 53 & 54
	118511	N75620	Hs.43157	ESTs	Seq ID 54 & 55
	101447	M21305		gb:Human alpha satellite and satellite 3	Seq ID 56 & 57
45	314941	AA515902	Hs.130650	ESTs .	Seq ID 58 & 59
	332466	AB018259	Hs.118140	KIAA0716 gene product	Seq ID 60 & 61
	313513	AW298600	Hs.141840	ESTs, Weakly similar to S59501 interfero	Seq ID 62 & 63
	313556	AA628517	Hs.118502	ESTs	Seq ID 64 & 65
50	313665	AW751201	Hs.51233	ESTs	Seq ID 66 & 67
50	314372	AL040178	Hs.142003	ESTs G pertoin equaled supporter 51	Seq ID 68 & 69 Seq ID 70 & 71
	429276 101345	AF056085 NM_005795	Hs.198612 Hs.152175	G protein-coupled receptor 51 calcitonin receptor-like	Seq ID 72 & 73
	418994	AA296520	Hs.89546	selectin E (endothelial adhesion molecul	Seq ID 74 & 75
	103850	AA187101	Hs.213194	hypothetical protein MGC10895	Seq ID 76 & 77
55	133260	AA403045	Hs.6906	Homo sapiens cONA: FLJ23197 fis, clone R	Seq ID 78 & 79
	101097	BE245301	Hs.89414	chemokine (C-X-C motif), receptor 4 (fus	Seq ID 80 & 81
	104786	AA027167	Hs.10031	KIAA0955 protein	Seq 1D 82 & 83
	132173	X89426	Hs.41716	endothelial cell-specific molecule 1	Seq ID 84 & 85
60	100420	D86983	Hs.118893	Metanoma associated gene	Seq ID 86 & 87
60	111018	A1287912	Hs.3628	mitogen-activated protein kinase kinase	Seq ID 88 & 89
	108507	AI554545	Hs.68301	ESTS  shortholiness A2 amus IVC (adoselis	Seq ID 90 & 91 Seq ID 92 & 93
	104894 118511	AF065214 N75620	Hs.18858 Hs.43157	phospholipase A2, group IVC (cytosolic, ESTs	Seg ID 94 & 95
	125609	AA868063	Hs.104576	carbohydrate (keratan sulfate Gal-6) sul	Seq ID 96 & 97
65	101543	M31166	Hs.2050	pentaxin-related gene, rapidly induced b	Seq ID 98 & 99
••	102241	NM_007351	Hs.268107	multimerin	Seq ID 100 & 101
	101560	AW958272	Hs.347326	intercellular adhesion molecule 2	Seq ID 102 & 103
	103280	U84722	Hs.76206	cadherin 5, type 2, VE-cadherin (vascula	Seq ID 104 & 105
70	105826	AA478756	Hs.194477	E3 ubiquitin ligase SMURF2	Seq ID 106 & 107
70	102804	NM_002318	Hs.83354	lysyl oxidase-like 2	Seq ID 108 & 109
	131647	AA359615	Hs.30089	ESTs	Seq ID 110 & 111
	103095	NM_005424	Hs.78824	tyrosine kinase with immunoglobulin and placental growth factor, vascular endoth	Seq ID 112 & 113
	103037 100405	BE018302 AW291587	Hs.2894 Hs.82733	placental grown factor, vascular endoth nidogen 2	Seq ID 114 & 115 Seq ID 116 & 117
75	102012	BE259035	Hs.118400	singed (Drosophila)-like (sea urchin fas	Seq ID 118 & 119
, 5	102012	<u> </u>	113.77000	origina (procedums/men food engine rea	004 ID 110 W 110

	101261	D30857	Hs.82353	protein C receptor, endothelial (EPCR)	Seq ID 120 & 121
	105729	H46612	Hs.293815	Homo sapiens HSPC285 mRNA, partial cds	Seq ID 122 & 123
	107216	D51069	Hs.211579	melanoma cell adhesion molecule	Seq ID 124 & 125
	131080	NM_001955	Hs.2271	endothelin 1	Seq ID 126 & 127
5	131486	F06972	Hs.27372	BMX non-receptor tyrosine kinase	Seq ID 128 & 129
•	134299	AW580939	Hs.97199	complement component C1q receptor	Seq ID 130 & 131
	134983	D28235	Hs.196384	prostaglandin-endoperoxide synthase 2 (p	Seq ID 132 & 133
	115827	AA428000	Hs.283072	actin related protein 2/3 complex, subun	Seq ID 134 & 135
10	133614	NM_003003	Hs.75232	SEC14 (S. cerevisiae)-like 1	Seq ID 136 & 137
10	116483	AJ346201	Hs.76118	ubiquitin carboxyl-terminal esterase L1	Seq ID 138 & 139
	132546	M24283	Hs.168383	intercellular adhesion molecule 1 (CD54)	Seq ID 140 & 141
	133678	AW247252	NA	nucleoside phosphorylase	Seq ID 142 & 143
	130184	H58306	Hs.15165	retinoic acid induced 14	Seq ID 144 & 145
	134786	T29618	Hs.89640	TEK tyrosine kinase, endothelial (venous	Seq ID 146 & 147
15	129371	X06828	Hs.110802	von Willebrand factor	Seq ID 148 & 149
	418506	AA084248	Hs.85339	G protein-coupled receptor 39	Seq ID 150 & 151
	322262	AA632012	Hs.188746	ESTs	Seq ID 152 & 153
	312173	Al821409	Hs.304471	EST ·	Seq ID 154 & 155
	319795	AB037821	Hs.146858	protocadherin 10	Seq ID 156 & 157
20			Hs.13957	ESTs	
20	313978	AI870175		· ·	Seq ID 158 & 159
	306840	A1077477	Hs.307912	ESTs	Seq ID 160 & 161
	310272	AF216389	Hs.148932	sema domain, transmembrane domain (TM),	Seq ID 162 & 163
	310272	AF216389	Hs.148932	sema domain, transmembrane domain (TM),	Seq ID 164 & 165
~ -	315044	BE547674	Hs.204169	ESTs, Weakly similar to S65657 alpha-1C-	Seq ID 166 & 167
25	321325	AB033100	Hs.300646	KIAA1274 protein (similar to mouse palad	Seq ID 168 & 169
	321325	AB033100	Hs.300646	KIAA1274 protein (similar to mouse palad	Seq ID 170 & 171
	303251	AF240635	Hs.115897	protocadherin 12	Seq ID 172 & 173
	302378	AL109712	Hs.296506	Homo sapiens mRNA full length insert cDN	Seq ID 174 & 175
	319267	F11802	Hs.6818	ESTs	Seq ID 176 & 177
30	310442	AW072215	Hs.208470	ESTs	Seq ID 178 & 179
-	300469	BE301708	Hs.233955	hypothetical protein FLJ20401	Seq ID 180 & 181
	331237	W87874	Hs.25277	Homo sapiens cDNA FLJ10717 fis; clone NT	Seq ID 182 & 183
	330968		Hs.23748	ESTs	Seq ID 184 & 185
		R44557			
35	301822	X17033	Hs.271986	integrin, atpha 2 (CD49B, alpha 2 subuni	Seq ID 186 & 187
33	422573	AW297985	Hs.295726	Integrin, alpha V (vitronectin receptor	Seq ID 188 & 189
	133061	Al186431	Hs.296638	prostate differentiation factor	Seq ID 190 & 191
	135235	AW298244	Hs.266195	ESTs	Seq ID 192 & 193
	101192	BE247295	Hs.78452	solute carrier family 20 (phosphate tran	Seq ID 194 & 195
	113195	H83265	Hs.8881	ESTs, Weakly similar to S41044 chromosom	Seq ID 196 & 197
40	101741	NM_003199	Hs.326198	transcription factor 4	Seq ID 198 & 199
	321911	AF026944	Hs.293797	ESTs	Seq ID 200 & 201
	320635	N50617	Hs.80506	small nuclear ribonucleoprotein polypept	Seq ID 202 & 203
	326230			NM_017643:Homo saplens hypothetical prot	Seg ID 204 & 205
	132968	AF234532	Hs.61638	myosin X	Seq ID 206 & 207
45	135073	W55956	Hs.94030	Homo sapiens mRNA; cDNA DKFZp586E1624 (f	Seq ID 208 & 209
10	108937	AL050107	Hs.24341	transcriptional co-activator with PDZ-bi	Seq ID 210 & 211
	116430	AK001531	Hs.66048	hypothetical protein FLI10669	Seq ID 212 & 213
				Hypothetical protein Ft. 10005	
	104877	Al138635	Hs.22968	Homo sapiens clone IMAGE:451939, mRNA se	Seq ID 214 & 215
50	122697	AA420683	Hs.98321	hypothetical protein FLJ14103	Seq ID 216 & 217
50	112522	R68857	Hs.265499	ESTs	Seq ID 218 & 219
	304782	AA582081		gb:nn32h08.s1 NCI_CGAP_Gas1 Homo sapiens	Seq ID 220 & 221
	312802	AA644669	Hs.193042	ESTs	Seq ID 222 & 223
	302680	AW192334	Hs.38218	ESTs	Seq ID 224 & 225
	326198			Phase 2 & 3 Exons	Seq ID 226 & 227
55	331019	NM_006033	Hs.65370	lipase; endothelial	Seq ID 228 & 229

#### TABLE 8

Seq ID NO: 1 DNA sequence

75

5 Nucleic Acid Accession #: NM_001400 244-2208 (underlined sequences correspond to start and stop codons)) Coding sequence: 10 11 21 31 41 51 GTCGGGGGCA GCAGCAAGAT GCGAAGCGAG CCGTACAGAT CCCGGGCTCT CCGAACGCAA 60 CTTCGCCCTG CTTGAGCGAG GCTGCGGTTT CCGAGGCCCT CTCCAGCCAA GGAAAAGCTA CACAAAAAGC CTGGATCACT CATCGAACCA CCCCTGAAGC CAGTGAAGGC TCTCTCGCCT 180 15 CGCCCTCTAG CGTTCGTCTG GAGTAGCGCC ACCCCGGCTT CCTGGGGACA CAGGGTTGGC 240 ACCATGGGGC CCACCAGCGT CCCGCTGGTC AAGGCCCACC GCAGCTCGGT CTCTGACTAC 300 GTCAACTATG ATATCATCGT CCGGCATTAC AACTACACGG GAAAGCTGAA TATCAGCGCG 360 GACAAGGAGA ACAGCATTAA ACTGACCTCG GTGGTGTTCA TTCTCATCTG CTGCTTTATC 420 ATCCTGGAGA ACATCTTTGT CTTGCTGACC ATTTGGAAAA CCAAGAAATT CCACCGACCC 480 20 ATGTACTATT TTATTGGCAA TCTGGCCCTC TCAGACCTGT TGGCAGGAGT AGCCTACACA GCTAACCTGC TCTTGTCTGG GGCCACCACC TACAAGCTCA CTCCCGCCCA GTGGTTTCTG 600 CGGGAAGGGA GTATGTTTGT GGCCCTGTCA GCCTCCGTGT TCAGTCTCCT CGCCATCGCC 660 ATTGAGCGCT ATATCACAAT GCTGAAAATG AAACTCCACA ACGGGAGCAA TAACTTCCGC 720 CTCTTCCTGC TAATCAGCGC CTGCTGGGTC ATCTCCCTCA TCCTGGGTGG CCTGCCTATC 780 25 ATGGGCTGGA ACTGCATCAG TGCGCTGTCC AGCTGCTCCA CCGTGCTGCC GCTCTACCAC 840 AAGCACTATA TCCTCTTCTG CACCACGGTC TTCACTCTGC TTCTGCTCTC CATCGTCATT 900 CTGTACTGCA GAATCTACTC CTTGGTCAGG ACTCGGAGGCC GCCGCCTGAC GTTCCGCAAG AACATTTCCA AGGCCAGCCG CAGCTCTGAG AAGTCGCTGG CGCTGCTCAA GACCGTAATT 1020 ATCGTCCTGA GCGTCTTCAT CGCCTGCTGG GCACCGCTCT TCATCCTGCT CCTGCTGGAT 1080 30 GTGGGCTGCA AGGTGAAGAC CTGTGACATC CTCTTCAGAG CGGAGTACTT CCTGGTGTTA 1140 GCTGTGCTCA ACTCCGGCAC CAACCCCATC ATTTACACTC TGACCAACAA GGAGATGCGT 1200 CGGGCCTTCA TCCGGATCAT GTCCTGCTGC AAGTGCCCGA GCGGAGACTC TGCTGGCAAA 1260 TTCAAGCGAC CCATCATCGC CGGCATGGAA TTCAGCCGCA GCAAATCGGA CAATTCCTCC 1320 CACCCCAGA AAGACGAAGG GGACAACCCA GAGACCATTA TGTCTTCTGG AAACGTCAAC 1380 35 TCTTCTTCCT AGAACTGGAA GCTGTCCACC CACCGGAAGC GCTCTTTACT TGGTCGCTGG CCACCCCAGT GTTTGGAAAA AAATCTCTGG GCTTCGACTG CTGCCAGGGA GGAGCTGCTG CAAGCCAGAG GGAGGAAGGG GGAGAATACG AACAGCCTGG TGGTGTCGGG TGTTGGTGGG 1560 TAGAGTTAGT TCCTGTGAAC AATGCACTGG GAAGGGTGGA GATCAGGTCC CGGCCTGGAA 1620 TATATATTCT ACCCCCTGG AGCTTTGATT TTGCACTGAG CCAAAGGTCT AGCATTGTCA 1680 40 AGCTCCTAAA GGGTTCATTT GGCCCCTCCT CAAAGACTAA TGTCCCCATG TGAAAGCGTC 1740 TCTTTGTCTG GAGCTTTGAG GAGATGTTTT CCTTCACTTT AGTTTCAAAC CCAAGTGAGT GTGTGCACTT CTGCTTCTTT AGGGATGCCC TGTACATCCC ACACCCCACC CTCCCTTCCC 1860 TTCATACCCC TCCTCAACGT TCTTTTACTT TATACTTTAA CTACCTGAGA GTTATCAGAG 1920 CTGGGGTTGT GGAATGATCO ATCATCTATA GCAAATAGGC TATGTTGAGT ACGTAGGCTG
TGGGAAGATG AAGATGGTTT GGAGGTGTAA AACAATGTCC TTCGCTGAGG CCAAAGTTTC 1980 45 2040 CATGTAAGCG GGATCCGTTT TTTGGAATTT GGTTGAAGTC ACTTTGATTT CTTTAAAAAA 2100 CATCTITTCA ATGAAATGTG TTACCATTTC ATATCCATTG AAGCCGAAAT CTGCATAAGG 2160 AAGCCCACTT TATCTAAATG ATATTAGCCA GGATCCTTGG TGTCC<u>TAG</u>GA GAAACAGACA 2220 AGCAAAACAA AGTGAAAACC GAATGGATTA ACTTTTGCAA ACCAAGGGAG ATTTCTTAGC 2280 50 AAATGAGTCT AACAAATATG ACATCCGTCT TTCCCACTTT TGTTGATGTT TATTTCAGAA 2340 TCTTGTGTGA TTCATTTCAA GCAACAACAT GTTGTATTTT GTTGTGTTAA AAGTACTTTT 2400 CTTGATTTTT GAATGTATTT GTTTCAGGAA GAAGTCATTT TATGGATTTT TCTAACCCGT 2460 GTTAACTTTT CTAGAATCCA CCCTCTTGTG CCCTTAAGCA TTACTTTAAC TGGTAGGGAA 2520 CGCCAGAACT TTTAAGTCCA GCTATTCATT AGATAGTAAT TGAAGATATG TATAAATATT 2580 55 ACAAAGAATA AAAATATATT ACTGTCTCTT TAGTATGGTT TTCAGTGCAA TTAAACCGAG 2640 AGATGTCTTG TTTTTTTAAA AAGAATAGTA TTTAATAGGT TTCTGACTTT TGTGGATCAT 2700 TTTGCACATA GCTTTATCAA CTTTTAAACA TTAATAAACT GATTTTTTTA AAG 60 Seq ID NO: 2 Protein sequence: Protein Accession #: NP_001391 65 21 31 11 41 51 MGPTSVPLVK AHRSSVSDYV NYDIIVRHYN YTGKLNISAD KENSIKLTSV VPILICCFII 60 LENIFULLTI WKTKKFHRPM YYPIGNLALS DLLAGVAYTA NLLLSGATTY KLTPAQWFLR 120 EGSMFVALSA SVFSLLAIAI ERYITMLKMK LHNGSNNFRL PLLISACWVI SLILGGLPIM 180 70 GWNCISALSS CSTVLPLYHK HYILFCTTVF TLLLLSIVIL YCRIYSLVRT RSRRLTFRKN 240 ISKASRSSEK SLALLKTVII VLSVFIACNA PLFILLLLDV GCKVKTCDIL FRABYFLVLA VLNSGTNPII YTLTNKEMRR APIRIMSCCK CPSGDSAGKF KRPIIAGMEF SRSKSDNSSH POKDEGDNPE TIMSSGNVNS SS

5 Seq ID NO: 3 Nucleotide sequence: 10 Nucleic Acid Accession #: NM 016242 Coding sequence: 79-864 (underlined sequences correspond to start and stop codons)) 21 31 41 51 15 AAGGCCCTGC CAGCTTGGGA GGGAATTGTC CCTGCCTGCT TCTGGAGAAA GAAGATATTG 60 ACACCATCTA CGGGCACCAT GGAACTGCTT CAAGTGACCA TTCTTTTTCT TCTGCCCAGT ATTTGCAGCA GTAACAGCAC AGGTGTTTTA GAGGCAGCTA ATAATTCACT TGTTGTTACT 120 180 ACAACAAAAC CATCTATAAC AACACCAAAC ACAGAATCAT TACAGAAAAA TGTTGTCACA 240 20 CCAACAACTG GAACAACTCC TAAAGGAACA ATCACCAATG AATTACTTAA AATGTCTCTG 300 ATGTCAACAG CTACTTTTTT AACAAGTAAA GATGAAGGAT TGAAAGCCAC AACCACTGAT 360 GTCAGGAAGA ATGACTCCAT CATTTCAAAC GTAACAGTAA CAAGTGTTAC ACTTCCCAAT 420 GCTGTTTCAA CATTACAAAG TTCCAAACCC AAGACTGAAA CTCAGAGTTC AATTAAAACA 480 ACAGAAATAC CAGGTAGTGT TCTACAACCA GATGCATCAC CTTCTAAAAC TGGTACATTA 25 ACCTCAATAC CAGTTACAAT TCCAGAAAAC ACCTCACAGT CTCAAGTAAT AGACACTGAG 600 GGTGGAAAAA ATGCAAGCAC TTCAGCAACC AGCCGGTCTT ATTCCAGTAT TATTTTGCCG 660 GTGGTTATTG CTTTGATTGT AATAACACTT TCAGTATTTG TTCTGGTGGG TTTGTACCGA 720 ATGTGCTGGA AGGCAGATCC GGGCACACCA GAAAATGGAA ATGATCAACC TCAGTCTGAT 780 AAAGAGAGCG TGAAGCTTCT TACCGTTAAG ACAATTTCTC ATGAGTCTGG TGAGCACTCT 840 30 GCACAAGGAA AAACCAAGAA C<u>TGA</u>CAGCTT GAGGAATTCT CTCCACACCT AGGCAATAAT 900 TACGCTTAAT CTTCAGCTTC TATGCACCAA GCGTGGAAAA GGAGAAAGTC CTGCAGAATC AATCCCGACT TCCATACCTG CTGCTGG 35 Seq ID NO: 4 Protein sequence: Protein Accession #: NP_057326 31 40 MELLQVTILF LLPSICSSNS TGVLEAANNS LVVTTTKPSI TTPNTESLQK NVVTPTTGTT 60 PKGTITNELL KMSLMSTATF LTSKDEGLKA TTTDVRKNDS IISNVTVTSV TLPNAVSTLQ 120 SSKPKTETQS SIKTTEIPGS VLQPDASPSK TGTLTSIPVT IPENTSQSQV IDTEGGKNAS 180 TSATSRSYSS IILPVVIALI VITLSVFVLV GLYRMCWKAD PGTPENGNDQ PQSDKESVKL 240 45 LTVKTISHES GEHSAQGKTK N Seq ID NO: 5 Nucleotide sequence: Nucleic Acid Accession #: NM_002205 , 50 24..3173 (underlined sequences correspond to start and stop codons) Coding sequence: 21 31 11 CAGGACAGGG AAGAGCGGGC GCTATGGGGA GCCGGACGCC AGAGTCCCCT CTCCACGCCG 55 60 TGCAGCTGCG CTGGGGCCCC CGGCGCCGAC CCCCGCTCGT GCCGCTGCTG TTGCTGCTCG 120 TGCCGCCGCC ACCCAGGGTC GGGGGCTTCA ACTTAGACGC GGAGGCCCCA GCAGTACTCT 180 CGGGGCCCCC GGGCTCCTTC TTCGGATTCT CAGTGGAGTT TTACCGGCCG GGAACAGACG 240 300 GGGTCAGTGT GCTGGTGGGA GCACCCAAGG CTAATACCAG CCAGCCAGGA GTGCTGCAGG 60 GTGGTGCTGT CTACCTCTGT CCTTGGGGTG CCAGCCCCAC ACAGTGCACC CCCATTGAAT 360 TTGACAGCAA AGGCTCTCGG CTCCTGGAGT CCTCACTGTC CAGCTCAGAG GGAGAGGAGC CTGTGGAGTA CAAGTCCTTG CAGTGGTTCG GGGCAACAGT TCGAGCCCAT GGCTCCTCCA 480 TCTTGGCATG CGCTCCACTG TACAGCTGGC GCACAGAGAA GGAGCCACTG AGCGACCCCG 540 TGGGCACCTG CTACCTCTCC ACAGATAACT TCACCCGAAT TCTGGAGTAT GCACCCTGCC 600 65 GCTCAGATTT CAGCTGGGCA GCAGGACAGG GTTACTGCCA AGGAGGCTTC AGTGCCGAGT 660 TCACCAAGAC TGGCCGTGTG GTTTTAGGTG GACCAGGAAG CTATTTCTGG CAAGGCCAGA 720 TCCTGTCTGC CACTCAGGAG CAGATTGCAG AATCTTATTA CCCCGAGTAC CTGATCAACC 780 TGGTTCAGGG GCAGCTGCAG ACTCGCCAGG CCAGTTCCAT CTATGATGAC AGCTACCTAG 840 GATACTCTGT GGCTGTTGGT GAATTCAGTG GTGATGACAC AGAAGACTTT GTTGCTGGTG 900 TGCCCAAAGG GAACCTCACT TACGGCTATG TCACCATCCT TAATGGCTCA GACATTCGAT 960 CCCTCTACAA CTTCTCAGGG GAACAGATGG CCTCCTACTT TGGCTATGCA GTGGCCGCCA 1020 CAGACGTCAA TGGGGACGGG CTGGATGACT TGCTGGTGGG GGCACCCCTG CTCATGGATC 1080 GGACCCCTGA CGGGCGGCCT CAGGAGGTGG GCAGGGTCTA CGTCTACCTG CAGCACCCAG 1140 CCGGCATAGA GCCCACGCCC ACCCTTACCC TCACTGGCCA TGATGAGTTT GGCCGATTTG 1200 GCAGCTCCTT GACCCCCCTG GGGGACCTGG ACCAGGATGG CTACAATGAT GTGGCCATCG

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21

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Seq ID NO: 21 <u>Nucleotide sequence:</u>
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5	AAGCAGTACT	TTCCDDTDCC	<b>ልጥጥ</b> ጉልር ልጥ	GAAACGGGAC	שריים אם הדום. ביים ממחדים מ	CLLALCECALE	600
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      MPALHIEDLP EKEKLKMEVE QLRKEVKLQR QQVSKCSEEI KNYIEERSGE DPLVKGIPED
      KNPFKEKGSC VIS
      Seq ID NO: 49 Nucleotide sequence:
75
      Nucleic Acid Accession #: XM_051896
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Coding sequence: 139..2388 (underlined sequences correspond to start and stop codons)

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      GAAACCAAAA TGTCATTTAT AGATCCTTAC CAGCACATTA TAGTGGAGCA CCAGTATTCC
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      CACAAGTTTA CGGTAGTGGT GTTACGTGCC ACCAAAGTGA CAAAGGGGGC CTTTGGTGAC
      ATGCTTGATA CTCCAGATCC CTATGTGGAA CTTTTTATCT CTACAACCCC TGACAGCAGG
                                                                          240
      AAGAGAACAA GACATTTCAA TAATGACATA AACCCTGTGT GGAATGAGAC CTTTGAATTT
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      ATTTTGGATC CTAATCAGGA AAATGTTTTG GAGATTACGT TAATGGATGC CAATTATGTC
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      ATGGATGAAA CTCTAGGGAC AGCAACATTT ACTGTATCTT CTATGAAGGT GGGAGAAAAG
                                                                          420
      AAAGAAGTTC CTTTTATTTT CAACCAAGTC ACTGAAATGG TTCTAGAAAT GTCTCTTGAA
      GTTTGCTCAT GCCCAGACCT ACGATTTAGT ATGGCTCTGT GTGATCAGGA GAAGACTTTC
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      AGACAACAGA GAAAAGAACA CATAAGGGAG AGCATGAAGA AACTCTTGGG TCCAAAGAAT
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      AGTGAAGGAT TGCATTCTGC ACGTGATGTG CCTGTGGTAG CCATATTGGG TTCAGGTGGG
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      GGTTTCCGAG CCATGGTGGG ATTCTCTGGT GTGATGAAGG CATTATACGA ATCAGGAATT
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      CTGGATTGTG CTACCTACGT TGCTGGTCTT TCTGGCTCCA CCTGGTATAT GTCAACCTTG
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      TATTCTCACC CTGATTTTCC AGAGAAAGGG CCAGAGGAGA TTAATGAAGA ACTAATGAAA
                                                                          840
      AATGTTAGCC ACAATCCCCT TTTACTTCTC ACACCACAGA AAGTTAAAAG ATATGTTGAG
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      TCTTTATGGA AGAAGAAAG CTCTGGACAA CCTGTCACCT TTACTGATAT CTTTGGGATG
      TTAATAGGAG AAACACTAAT TCATAATAGA ATGAATACTA CTCTGAGCAG TTTGAAGGAA
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      ANAGTTAATA CTGCACAATG CCCTTTACCT CTTTTCACCT GTCTTCATGT CANACCTGAC
                                                                        1080
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      GTTAAGAAGT ATGAAGAAAA CCCCTTGCAT TTCTTAATGG GTGTCTGGGG CAGTGCCTTT
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      GAGGAAGAAT TAGAAAATAT TACCACAAAG CATATTGTGA GTAATGATAG CTCGGACAGT
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      CTGGATGTCA AAAGTAAAAA GATTCATGTA GTGGACAGTG GGCTCACATT TAACCTGCCG
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                                                                        1980
      ATCCACTTG TTCTGGCCAA CATCAACTTC AGAAAGTACA GGGCTCCAGG TGTTCCAAGG
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                                                                        2160
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      TATAGAAGAC AGAATCCATC TCGTTGCTCT GTTTCCCTTA GTAATGTTGA GGCAAGAAGA
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      TTAGCTACAT TTTCAGTCAG TATGAACTTC CTGATACAAA TGTAGGGATA TATACTGTAT
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      TTTTAAACAT TTCTCACCAA CTTTCTTATG TGTGTTCTTT TTAAAAATTT TTTTTCTTTT
                                                                        2640
      AAAATATTTA ACAGTTCAAT CTCAATAAGA CCTCGCATTA TGTATGAATG TTATTCACTG
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      Seq ID NO: 50 Protein sequence:
      Protein Accession #: XP_051896
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                                                                          60
      RHFNNDINPV WNETFBFILD PNQENVLEIT LMDANYVMDE TLGTATFTVS SMKVGEKKEV
                                                                         120
60
      PFIFNQVTEM VLEMSLEVCS CPDLRFSMAL CDQEKTFRQQ RKEHIRESMK KLLGPKNSEG
                                                                         180
      LHSARDVPVV AILGSGGGFR AMVGFSGVMK ALYESGILDC ATYVAGLSGS TWYMSTLYSH
                                                                         240
      PDFPEKGPEE INEELMKNVS HNPLLLLTPQ KVKRYVESLW KKKSSGQPVT FTDIFGMLIG
                                                                         300
      ETLIHNRMNT TLSSLKEKVN TAQCPLPLFT CLHVKPDVSE LMFADWVBFS PYEIGMAKYG
      TFMAPDLIFGS KFFMGTVVKK YEENPLHFLM GVWGSAFSIL FNRVLGVSGS QSRGSTMEEE
                                                                         420
65
      LENITTKHIV SNDSSDSDDE SHEPKGTENE DAGSDYQSDN QASWIHRMIM ALVSDSALFN
                                                                         480
      TREGRAGKVH NFMLGLNINT SYPLSPLSDF ATODSFDDDE IDAAVADPDE FERIYEPLDV
                                                                         540
      KSKKIHVVDS GLTFNLPYPL ILRPQRGVDL IISFDFSARP SDSSPPFKEL LLAEKWAKMN
                                                                         600
      KLPFPKIDPY VFDREGLKEC YVFKPKNPDM EKDCPTIIHF VLANINFRKY KAPGVPRETE
                                                                         660
      EEKELADFDI FDDPESPFST FNFQYPNQAF KRLHDLMHFN TLNNIDVIKE AMVESIEYRR
70
      ONPSRCSVSL SNVBARRFFN KEFLSKPKA
      Seq ID NO: 51 Nucleotide sequence:
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Seq ID NO: 51 <u>Nucleotide sequence:</u>
Nucleic Acid Accession #: NM_006528
Coding sequence: 57..764 (underlined sequences correspond to start and stop codons)

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                                                                           120
      GCGATGCTGC TCAGGAGCCA ACAGGAAATA ACGCGGAGAT CTGTCTCCTG CCCCTAGACT
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      ACGGACCCTG CCGGGCCCTA CTTCTCCGTT ACTACTACGA CAGGTACACG CAGAGCTGCC
      GCCAGTTCCT GTACGGGGGC TGCGAGGGCA ACGCCAACAA TTTCTACACC TGGGAGGCTT
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      GCGACGATGC TTGCTGGAGG ATAGAAAAAG TTCCCAAAGT TTGCCGGCTG CAAGTGAGTG
                                                                           360
      TGGACGACCA GTGTGAGGGG TCCACAGAAA AGTATTTCTT TAATCTAAGT TCCATGACAT
                                                                           420
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      GTGAAAAATT CTTTTCCGGT GGGTGTCACC GGAACCGGAT TGAGAACAGG TTTCCAGATG
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      AAGCTACTTG TATGGGCTTC TGCGCACCAA AGAAAATTCC ATCATTTTGC TACAGTCCAA
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      AAGATGAGGG ACTGTGCTCT GCCAATGTGA CTCGCTATTA TTTTAATCCA AGATACAGAA
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      AGGATTGCAA ACGTGCATGT GCAAAAGCTT TGAAAAAGAA AAAGAAGATG CCAAAGCTTC
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      GCTTTGCCAG TAGAATCCGG AAAATTCGGA AGAAGCAATT <u>TTA</u>AACATTC TTAATATGTC
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      ATCITGTITG TCTTTATGGC TTATTTGCCT TTATGGTTGT ATCTGAAGAA TAATATGACA
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      GCATGAGGAA ACAAATCATT GGTGATTTAT TCACCAGTTT TTATTAATAC AAGTCACTTT
                                                                           900
      TTCAAAAATT TGGATTTTTT TATATATAAC TAGCTGCTAT TCAAATGTGA GTCTACCATT
                                                                           960
      TTTAATTTAT GGTTCAACTG TTTGTGAGAC GAATTCTTGC AATGCATAAG ATATAAAAGC
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      AAATATGACT CACTCATTTC TTGGGGTCGT ATTCCTGATT TCAGAAGAGG ATCATAACTG
                                                                          1080
      AAACAACATA AGACAATATA ATCATGTGCT TTTAACATAT TTGAGAATAA AAAGGACTAG
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      Protein Accession #: NP_006519
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      CROFLYGGCE GNANNFYTWE ACDDACWRIE KVPKVCRLOV SVDDQCEGST EKYFFNLSSM
                                                                           120
      TCEKFFSGGC HRNRIENRFP DEATCMGFCA PKKIPSFCYS PKDEGLCSAN VTRYYFNPRY
      RTCDAFTYTG CGGNDNNFVS REDCKRACAK ALKKKKMPK LRFASRIRKI RKKQF
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      Nucleic Acid Accession #: AA478778
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      ACTITITIAA AATGAATITI TITAAAACAA TCTAGCCATC ATCAAGGTGC TATAAGAGTT
                                                                           180
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      GTATAAAAGA TATTTTTGGC ATTTCTAGGC AAGTATCAGC CAATAAGTAT GTTAGTGATA
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      TCACAGATTG TACCAACTAT TAACTATGTT AAATAAGTAT TCAGTTTCAT GTGATCTCTG
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      GGAAAAAAT ATGCTGCCTT GGTGCTAATA TTGTATGTAT TTAAATGATC ATCTGACTCA
      GAAATATAAA CACTTTTAAT GAAAGGGAGG AACGGAAGGA CAATTTCCAG TGCACAGAAT
      CACTTGGATG AAATAAGACC AGCTCTTTAC CCTTATTTTT GGATATGCCT TTTTTGGAAG
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      AGACTTAGAC TTTATCCTTA TTGTTGTTAG TGTTGTTAAT ATTCGTTGCT TCAGCCCACG
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      GTGCCTTGGT CTCTCCACAA TCAAATGGAG GATCCCCCAA GCAGCTTCAT TACAGAGTGA
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      TATTGGGAAA GTGAGATCCT CTCACCATTT TGCCAAGATA CTCTAAAATG ACATCCAAGT
                                                                           660
      TTACCAGTAG AAAGACACAG GATGCACAGA ATGGGCATGA CCTTCAGCTC ACGAGCACAC
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                                                                           780
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      GCTGGTGATG TGACTTCTCT TCAGGCCATG AGCCTAACAY CCTGCCGGTT TTCATGCCCG
                                                                           840
      CTGCAGTAAT GGACGTTTGT GTGAAGAAAT GAACTGTGGA GTACAAAATG CTTTGAGTCT
      TTCCGATTGC TCATTAATTC ACTTTTTGT TACTTCTTTC CAAAATGGAA GTGCTGAAGC
                                                                           960
      CATGGTCTTT CTGCCCCTCC AAGCTGATGA AGGGAAGCCT TTGCCAATGG CCCATGGAAG
                                                                          1020
      ACACTTGGTT TGAGAAACCC TGCCCACTTC CAAAGACCAA AGAGATTAGG AAAAGCCTGG
                                                                          1080
60
      CAGTATTCTC CAACTCCAAA CAAGCTCTAG AGTGCTCCAG GAAAAGTTAT ATTCAGTATA
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      CACCACACCC AAAAAAAAAA AAAAAAAAAA AAAA
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                                                                           120
      TACGCCAACG ACGCCTTCCC AGAGGAATAC GTGCCCACTG TGTTTGACCA CTATGCAGTT
                                                                           180
75
      ACTGTGACTG TGGGAGGCAA GCAACACTTG CTCGGACTGT ATGACACCGC GGGACAGGAG
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      GACTACAACC AGCTGAGGCC ACTCTCCTAC CCCAACACGG ATGTGTTTTT GATCTGCTTC
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TCTGTCGTAA ACCCTGCCTC TTACCACAAT GTCCAGGAGG AATGGGTCCC CGAGCTCAAG
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      CCAAAAACCT TGGCCCGTTT GCTGTATATG AAAGAGAAAC CTCTCACTTA CGAGCATGGT
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      GTGAAGCTCG CAAAAGCGAT CGGAGCACAG TGCTACTTGG AATGTTCAGC TCTGACTCAG
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      AAAGGTCTCA AAGCGGTTTT TGATGAAGCA ATCCTCACCA TTTTCCACCC CAAGAAAAAG
      AAGAAACGCT GTTCTGAGGG TCACAGCTGC TGTTCAATTA TCTGA
      Seq ID NO: 55 Protein sequence:
10
      Protein Accession #: NP_065714
      MNCKEGTDSS CGCRGNDEKK MLKCVVVGDG AVGKTCLLMS YANDAFPEEY VPTVFDHYAV
                                                                           60
15
      TVTVGGKOHL LGLYDTAGOE DYNOLRPLSY PNTDVFLICF SVVNPASYHN VOEEWVPELK
                                                                          120
      DCMPHVPYVL IGTQIDLRDD PKTLARLLYM KEKPLTYEHG VKLAKAIGAQ CYLECSALTQ
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      KGLKAVFDEA ILTIPHPKKK KKRCSEGHSC CSII
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      Seq ID NO: 56 Nucleotide sequence:
      Nucleic Acid Accession #: fgenesh prediction
      Coding sequence: 1-546 (underlined sequences correspond to start and stop codons)
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      ATGCCCTTGG GCAGCTCCGC CCCTGTGGCT TTGCAGGGTA ATGCCCACTT CCCTGCTGCT
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      TTGGTGCAAA GGCTCCTGGG TGGAGCTCGA ACTGAAACTC GCTTTGTGCC CGCAGCCCTG
                                                                          180
30
      CAGCTCGCCG GTGCCCTCGA CCTGCCCGCT GGGTCCTGTG CCTTTGAAGA GAGCACTTGC
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      GGCTTTGACT CCGTGTTGGC CTCTCTGCCG TGGATTTTAA ATGAGGAAGG CCAGCAACCT
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      TTCTGGTCCT CAGGAGACAT GTCTGACTGG GACTACTGGG TTGGCTGGCG GAAGTTAATT
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      CATTCTCCTC TGAGCACTCC AGGGTGGAGC AGGCAGGTTA GGCTCCAGTT GTTCCAGCTT
      CAGTTTGTCA AAGGCCAGAA CTTGGACGTA ACAGTGTACT GCAGGCTCCA GGGCAGTGAG
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      AAACCCTTTG AAACTGGTTC CATGGTTCCA TTCACCTTCA TGTACTGGAT CCACCATGGA
                                                                          540
      AAGTAG
      Seq ID NO: 57 Protein sequence:
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      Protein Accession #: fgenesh prediction
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      MALGSSAPVA LQGNAHFPAA FMAGIKCLWL FOVVPLGLPB LVORLLGGAR TETRFVPAAL
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45
      QLAGALDLPA GSCAPEESTC GFDSVLASLP WILNEEGQQP FWSSGDMSDW DYWVGWRKLI
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      HSPLSTPGWS RQVRLQLFQL QFVKGQNLDV TVYCRLQGSE KPPETGSMVP FTFMYWIHHG
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      Nucleic Acid Accession #: XM 050478
      Coding sequence: 27..4508 (underlined sequences correspond to start and stop codons)
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      GTGAGCCGCT CACAGTGTCT AAGATTGAAG ATGGAGGCAA GGCAGCTTTG TCCCAGAAGA
      TGAGGACTGG TGATGAGCTG GTGAATATCA ATGGCACTCC ATTATATGGC TCCCGCCAAG
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      AGGCCCTCAT TCTCATCAAA GGCTCCTTCC GGATTCTCAA GCTGATTGTC AGGAGGAGGA
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      ACGCCCCTGT CAGTAGGCCG CACTCATGGC ATGTGGCCAA GCTGCTGGAG GGATGCCCTG
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      AAGCAGCCAC CACCATGCAT TTCCCTTCTG AAGCCTTCAG CTTGTCCTGG CATTCTGGCT
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      CCTTCTCGGC CAGCTCAAAT GCTTCTGACT GTGCCCTTTC CCTCAGGCCA GAGGAGCCAG
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      CCTCTACAGA CTGCATCATG CAAGGCCCAG GGCCAACTAA GGCCCCCAGT GGCCGGCCTA
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			GAGGCTGATG				1560
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23			CATAATCCTC				2880
•			CCCAGGAAGC				2940
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			TTTCCTCCTC				3360
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	CAGAGGAGGA	GGAAGAGGAG	CTGCCACCCC	AGTATTTCAG	TTCAGAAACC	TCTGGTTCCT	3540
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			CAAAGTGTCC				3660
40			ATAAGGGGTC				3720
40			GGTGGGCTTT				3780
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			TTGCTGTCAC				4200
			GAGGCCAACC				4260
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			TCTGCCACAG				4560
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			TCCCAGCAGC				
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65						TTGTTTCATC	5220
65			GTTACCGCAG				5280
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			GGCGAGAAGA				5460
						TTCTTGCTCT	5520
70 .			CCTGCCTGGA				
, ,							5580
						TCCGGCCGGG	5640
						GGATCACGAG	5700
			CTGGCTAACA				5760
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      NEGDPTETLR QCFDDFADLV PFDSWEPLMR KLGLMDNEIK VAKABAAGHR DTLYTMLIKW
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                                                                          540
      NVTEYMSTLH ENIKKQSLMM LQMFEDLHIQ ESKINNLTVS LEMEKESLRG ECEDMLSKCR
                                                                          600
      NDFKFQLKDT BENLHVLNQT LAEVLFPMDN KMDKMSEQLN DLTYDMEILQ PLLEQGASLR
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      QTMTYEQPKE AIVIRKKIEN LTSAVNSLNF IIKELTKRHN LLRNEVQGRD DALERRINEY
                                                                          720
      ALEMEDGLNK TMTIINNAID FIODNYALKE TLSTIKDNSE IHHKCTSDME TILTFIPOFH
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      RLNDSIQTLV NDNQRYNFVL QVAKTLAGIP RDEKLNQSNF QKMYQMFNET TSQVRKYQQN
                                                                          840
      MSHLEEKLLL TTKISKNFET RLQDIESKVT QTLIPYYISV KKGSVVTNER DQALQLQVLN
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      SRFKALEAKS IHLSINFFSL NKTLHEVLTM CHNASTSVSE LNATIPKWIK HSLPDIQLLQ
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      KGLTEFVEPI IQIKTQAALS NSTCCIDRSL PGSLANVVKS QKQVKSLPKK INALKKPTVN
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      LTTVLIGRTQ RNTDNIIYPE EYSSCSRHPC QNGGTCINGR TSFTCACRHP FTGDNCTIKL
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      VEENALAPDF SKGSYRYAPM VAFFASHTYG MTIPGPILFN NLDVNYGASY TPRTGKFRIP
55
      YLGVYVFKYT IESFSAHISG FLVVDGIDKL AFRSENINSE IHCDRVLTGD ALLELNYGQE
      VWLRLAKGTI PAKFPPVTTF SGYLLYRT
      Seq ID NO: 102 DNA sequence
      Nucleic Acid Accession #: NM_000873.2
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      CCTCTTCGG TTACAGGACC CTGACTGTGG CCCTCTTCAC CCTGATCTGC TGTCCAGGAT
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      CGGATGAGAA GGTATTCGAG GTACACGTGA GGCCAAAGAA GCTGGCGGTT GAGCCCAAAG
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      GGTCCCTCGA GGTCAACTGC AGCACCACCT GTAACCAGCC TGAAGTGGGT GGTCTGGAGA
                                                                          240
      CCTCTCTAAA TAAGATTCTG CTGGACGAAC AGGCTCAGTG GAAACATTAC TTGGTCTCAA
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70
      ACATCTCCCA TGACACGGTC CTCCAATGCC ACTTCACCTG CTCCGGGAAG CAGGAGTCAA
      TGAATTCCAA CGTCAGCGTG TACCAGCCTC CAAGGCAGGT CATCCTGACA CTGCAACCCA
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      CTTTGGTGGC TGTGGGCAAG TCCTTCACCA TTGAGTGCAG GGTGCCCACC GTGGAGCCCC
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      TEGACAGCCT CACCCTCTTC CTGTTCCGTG GCAATGAGAC TCTGCACTAT GAGACCTTCG
                                                                          540
      GGAAGGCAGC CCCTGCTCCG CAGGAGGCCA CAGCCACATT CAACAGCACG GCTGACAGAG
                                                                          600
75
      AGGATGGCCA CCGCAACTTC TCCTGCCTGG CTGTGCTGGA CTTGATGTCT CGCGGTGGCA
                                                                          660
      ACATCTTCA CAAACACTCA GCCCGAAGA TGTTGGAGAT CTATGAGCCT GTGTCGGACA
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GCCAGATGGT CATCATAGTC ACGGTGGTGT CGGTGTTGCT GTCCCTGTTC GTGACATCTG
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       TGCGAGCGGC TTGGAGGAGG CTGCCCCAGG CCTTCCGGCC ATAGCAACCA TGAGTGGCAT
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       CAAACACCTG GACTT
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       Protein Accession #: NP_000864.1
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                                        31
       MSSPGYRTLT VALPTLICCP GSDEKVFEVH VRPKKLAVEP KGSLEVNCST TCNQPEVGGL
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15
       ETSLNKILLD EQAQWKHYLV SNISHDTVLQ CHFTCSGKQE SMNSNVSVYQ PPRQVILTLQ
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       PTLVAVGKSF TIECRVPTVE PLDSLTLFLF RGNETLHYET FGKAAPAPQE ATATFNSTAD
                                                                            180
       REDGHRNFSC LAVLDLMSRG GNIFHKHSAP KMLEIYEPVS DSQMVIIVTV VSVLLSLFVT
       SVLLCFIFGQ HLRQQRMGTY GVRAAWRRLP OAFRP
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       Nucleic Acid Accession #: NM_001795.2
       Coding sequence: 121-2475 (underlined sequences correspond to start and stop codons)
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      ATGCAGAGGC TCATGATGCT CCTCGCCACA TCGGGCGCCT GCCTGGGCCT GCTGGCAGTG
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       GCAGCAGTGG CAGCAGCAGG TGCTAACCCT GCCCAACGGG ACACCCACAG CCTGCTGCCC
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      ACCCACCGC GCCAAAAGAG AGATTGGATT TGGAACCAGA TGCACATTGA TGAAGAGAAA
      AACACCTCAC TTCCCCATCA TGTAGGCAAG ATCAAGTCAA GCGTGAGTCG CAAGAATGCC
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      AAGTACCTGC TCAAAGGAGA ATATGTGGGC AAGGTCTTCC GGGTCGATGC AGAGACAGGA
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      GACGTGTTCG CCATTGAGAG GCTGGACCGG GAGAATATCT CAGAGTACCA CCTCACTGCT
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      GTCATTGTGG ACAAGGACAC TGGTGAAAAC CTGGAGACTC CTTCCAGCTT CACCATCAAA
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      GTTCATGACG TGAACGACAA CTGGCCTGTG TTCACGCATC GGTTGTTCAA TGCGTCCGTG
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      CCTGAGTCGT CGGCTGTGGG GACCTCAGTC ATCTCTGTGA CAGCAGTGGA TGCAGACGAC
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      CCCACTGTGG GAGACCACGC CTCTGTCATG TACCAAATCC TGAAGGGGAA AGAGTATTTT
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      GCCATCGATA ATTCTGGACG TATTATCACA ATAACGAAAA GCTTGGACCG AGAGAAGCAG
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      GCCAGGTATG AGATCGTGGT GGAAGCGCGA GATGCCCAGG GCCTCCGGGG GGACTCGGGC
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      ACGGCCACCG TGCTGGTCAC TCTGCAAGAC ATCAATGACA ACTTCCCCTT CTTCACCCAG
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      ACCAGTACA CATTTGTCGT GCCTGAAGAC ACCCGTGTGG GCACCTCTGT GGGCTCTCTG
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      TTTGTTGAGG ACCCAGATGA GCCCCAGAAC CGGATGACCA AGTACAGCAT CTTGCGGGGC
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      GACTACCAGG ACGCTTTCAC CATTGAGACA AACCCCGCCC ACAACGAGGG CATCATCAAG
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      CCCATGAAGC CTCTGGATTA TGAATACATC CAGCAATACA GCTTCATCGT CGAGGCCACA
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      GACCCCACCA TCGACCTCCG ATACATGAGC CCTCCCGCGG GAAACAGAGC CCAGGTCATT
      ATCAACATCA CAGATGTGGA CGAGCCCCCC ATTTTCCAGC AGCCTTTCTA CCACTTCCAG
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      CTGAAGGAAA ACCAGAAGAA GCCTCTGATT GGCACAGTGC TGGCCATGGA CCCTGATGCG
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      GCTAGGCATA GCATTGGATA CTCCATCCGC AGGACCAGTG ACAAGGGCCA GTTCTTCCGA
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      GTCACAAAAA AGGGGGACAT TTACAATGAG AAAGAACTGG ACAGAGAAGT CTACCCCTGG
TATAACCTGA CTGTGGAGGC CAAAGAACTG GATTCCACTG GAACCCCCAC AGGAAAAGAA
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                                                                           1500
      TCCATTGTGC AAGTCCACAT TGAAGTTTTG GATGAGAATG ACAATGCCCC GGAGTTTGCC
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      AAGCCCTACC AGCCCAAAGT GTGTGAGAAC GCTGTCCATG GCCAGCTGGT CCTGCAGATC
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      GAGAACAACT TTACCCTCAC GGATAATCAC GATAACACGG CCAACATCAC AGTCAAGTAT
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      GGGCAGTTTG ACCGGGAGCA TACCAAGGTC CACTTCCTAC CCGTGGTCAT CTCAGACAAT
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      GGGATGCCAA GTCGCACGGG CACCAGCACG CTGACCGTGG CCGTGTGCAA GTGCAACGAG
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      CAGGGCGAGT TCACCTTCTG CGAGGATATG GCCGCCCAGG TGGGCGTGAG CATCCAGGCA
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      GTGGTAGCCA TCTTACTCTG CATCCTCACC ATCACAGTGA TCACCCTGCT CATCTTCCTG
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      CGGCGGCGC TCCGGAAGCA GGCCCGCGC CACGGCAAGA GCGTGCCGGA GATCCACGAG
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      CAGCTGGTCA CCTACGACGA GGAGGGCGGC GGCGAGATGG ACACCACCAG CTACGATGTG
      TOGGTGCTCA ACTOGGTGCG COGCGGCGGG GCCAAGCCCC CGCGGCCCGC GCTGGACGCC
                                                                           2160
      CGGCCTTCCC TCTATGCGCA GGTGCAGAAG CCACCGAGGC ACGCGCCTGG GGCACACGGA
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      GGGCCCGGGG AGATGGCAGC CATGATCGAG GTGAAGAAGG ACGAGGCGGA CCACGACGGC
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65
      GACGGCCCCC CCTACGACAC GCTGCACATC TACGGCTACG AGGGCTCCGA GTCCATAGCC
                                                                           2340
      GAGTCCCTCA GCTCCCTGGG CACCGACTCA TCCGACTCTG ACGTGGATTA CGACTTCCTT
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      AACGACTGGG GACCCAGGTT TAAGATGCTG GCTGAGCTGT ACGGCTCGGA CCCCCGGGAG
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      GAGCTGCTGT ATTAGGCGGC CGAGGTCACT CTGGGCCTGG GGACCCAAAC CCCCTGCAGC
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      GCACCCCTTC CTCGTGGGTC CCAGAGACCT CATCAGCCTT GGGATAGCAA ACTCCAGGTT
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      CCTGAAATAT CCAGGAATAT ATGTCAGTGA TGACTATTCT CAAATGCTGG CAAATCCAGG
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      CTGGTGTTCT-GTCTGGGCTC AGACATCCAC ATAACCCTGT CACCCACAGA CCGCCGTCTA
                                                                           2760
      ACTCAAAGAC TTCCTCTGGC TCCCCAAGGC TGCAAAGCAA AACAGACTGT GTTTAACTGC
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      TATCTGCCTG GAGGCAAAGG CCTGGACAGC TTGACTTGTG GGGCAGGATT CTCTGCAGCC
      CATTCCCAAG GGAGACTGAC CATCATGCCC TCTCTCGGGA GCCCTAGCCC TGCTCCAACT
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CCATACTCCA CTCCAAGTGC CCCACCACTC CCCAACCCCT CTCCAGGCCT GTCAAGAGGG
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      ATGCCAGTAA CTGTGCTGTA CTGAGCACTG AACCACATTC AGGGAAATGG CTTATTAAAC
                                                                          3180
      TTTGAAGCAA CTGTGAATTC ATTCTGGAGG GGCAGTGGAG ATCAGGAGTG ACAGATCACA
                                                                          3240
      GGGTGAGGGC CACCTCCACA CCCACCCCCT CTGGAGAAGG CCTGGAAGAG CTGAGACCTT
      GCTTTGAGAC TCCTCAGCAC CCCTCCAGTT TTGCCTGAGA AGGGGCAGAT GTTCCCGGAG
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      CAGAAGACGT CTCCCCTTCT CTGCCTCACC TGGTCGCCAA TCCATGCTCT CTTTCTTTTC
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      TCTGTCTACT CCTTATCCCT TGGTTTAGAG GAACCCAAGA TGTGGCCTTT AGCAAAACTG
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      CACCTTGGAG AAGTGGCATC AGTCAACAGA GAGGGGCAGG GAAGGAGACA CCAAGCTCAC
      CCTTCGTCAT GGACCGAGGT TCCCACTCTG GGCAAAGCCC CTCACACTGC AAGGGATTGT
      AGATAACACT GACTTGTTTG TTTTAACCAA TAACTAGCTT CTTATAATGA TTTTTTTACT
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      AATGATACTT ACAAGTTTCT AGCTCTCACA GACATATAGA ATAAGGGTTT TTGCATAATA
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                                                                          4020
      CTGTAACCTT CTATTTTCTA TAATTGTAGT AATTGCTCTA CAGATAATGT CTATATATTG
      GCCAAACTGG TGCATGACAA GTACTGTATT TITTTATACC TAAATAAAGA AAAATCTTTA
     GCCTGGGCAA CAAAAAA
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      Seq ID No: 105 Protein sequence:
      Protein Accession #: NP_001786.1
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      NTSLPHHVGK IKSSVSRKNA KYLLKGEYVG KVFRVDAETG DVFAIERLDR ENISEYHLTA
                                                                           120
      VIVDKDTGEN LETPSSFTIK VHDVNDNWPV FTHRLFNASV PESSAVGTSV ISVTAVDADD
                                                                           180
30
      PTVGDHASVM YQILKGKEYF AIDNSGRIIT ITKSLDREKQ ARYEIVVEAR DAQGLRGDSG
                                                                           240
      TATVLVTLQD INDNFPFFTQ TKYTFVVPED TRVGTSVGSL FVEDPDEPQN RMTKYSILRG
                                                                            300
      DYQDAFTIET NPAHNEGIIK PMKPLDYEYI QQYSFIVEAT DPTIDLRYMS PPAGNRAQVI
      INITDVDEPP IFQQPFYHFQ LKENQKKPLI GTVLAMDPDA ARHSIGYSIR RTSDKGQFFR
                                                                           420
      VTKKGDIYNE KELDREVYPW YNLTVEAKEL DSTGTPTGKE SIVQVHIEVL DENDNAPEPA
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35
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      GQFDREHTKV HFLPVVISDN GMPSRTGTST LTVAVCKCNE QGEFTFCEDM AAQVGVSIQA
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      VVAILLCILT ITVITLLIFL RRRLRKQARA HGKSVPEIHE QLVTYDEEGG GEMDTTSYDV
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      SVLNSVRRGG AKPPRPALDA RPSLYAQVQK PPRHAPGAHG GPGEMAAMIE VKKDBADHDG
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      Coding sequence: 1-474 (underlined sequences correspond to start and stop codons)
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      CTTGATCCAA AGTGGAATCA GCATTATGAC CTGTATATTG GAAAGTCTGA TTCAGTTACG
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      ATCAGTGTAT GGAATCACAA GAAGATCCAT AAGAAACAAG GTGCTGGATT TCTCGGTTGT
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      GTTCGTCTTC TTTCCAATGC CATCAACCGC CTCAAAGACA CTGGTTATCA GAGGTTGGAT
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      TTATGCAAAC TCGGGCCAAA TGACAATGAT ACAGTTAGAG GACAGATAGT AGTAAGTCTT
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      CAGTCCAGAG ACCGAATAGG CACAGGAGGA CAAGTTGTGG ACTGCAGTCG TTTATTTGAT
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      AACGATTTAC CAGACGGAGC TCATTATTTG TGGACTTGGA AAGATAGATG TTAATGACTG
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      GAAGGTAAAC ACCCGGTTAA AACACTGTAC ACCAGACAGC AACATTGTCA AATGGTTCTG
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      GAAAGCTGTG GAGTTTTTTG ATGAAGAGCG ACGAGCAAGA TTGCTTCAGT TTGTGACAGG
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      ATCCTCTCGA GTGCCTCTGC AGGGCTTCAA AGCATTGCAA GGTGCTGCAG GCCCGAGACT
                                                                           660
      CTTTACCATA CACCAGATTG ATGCCTGCAC TAACAACCTG CCGAAAGCCC ACACTTGCTT
                                                                            720
60
      CAATCGAATA GACATTCCAC CCTATGAAAG CTATGAAAAG CTATATGAAA AGCTGCTAAC
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      AGCCATTGAA GAAACATGTG GATTTGCTGT GGAATGACAA GCTTCAAGGA TTTACCCAGG
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      ISVWNHKKIH KKQGAGFLGC VRLLSNAINR LKDTGYQRLD LCKLGPNDND TVRGQIVVSL
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      QSRDRIGTGG QVVDCSRLFD NDLPDGAHYL WTWKDRC
      Seq ID NO: 108 DNA sequence
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      Nucleic Acid Accession #: NM_002318.1
      Coding sequence: 248-2572 (underlined sequences correspond to start and stop codons)
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51

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      GAGAGOGOGG TOCCAGCOCG GCCACTGOGG ATCCCTGAAA CCAAAAAGCT CCTGCTGCTT
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      CTGTACCCCG CCTGTCCCTC CCAGCTGCGC AGGGCCCCTT CGTGGGATCA TCAGCCCGAA
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      GACAGGGATG GAGAGGCCTC TGTGCTCCCA CCTCTGCAGC TGCCTGGCTA TGCTGGCCCT
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      CCTGTCCCCC CTGAGCCTGG CACAGTATGA CAGCTGGCCC CATTACCCCG AGTACTTCCA
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      GCAACOGGCT CCTGAGTATC ACCAGCCCCA GGCCCCCGCC AACGTGGCCA AGATTCAGCT
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      GCGCCTGGCT GGGCAGAAGA GGAAGCACAG CGAGGGCCGG GTGGAGGTGT ACTATGATGG
                                                                          480
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      CCTGAATATC CAGGTGGAGG ACATTCGGAT TCGAGCCATC CTCTCAACCT ACCGCAAGCG
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      CACCCCAGTG ATGGAGGGCT ACGTGGAGGT GAAGGGGGC AAGACCTGGA AGCAGATCTG
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      TGACAAGCAC TGGACGGCCA AGAATTCCCG CGTGGTCTGC GGCATGTTTG GCTTCCCTGG
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      GGAGAGGACA TACAATACCA AAGTGTACAA AATGTTTGCC TCACGGAGGA AGCAGCGCTA
                                                                        1020
      CTGGCCATTC TCCATGGACT GCACCGGCAC AGAGGCCCAC ATCTCCAGCT GCAAGCTGGG
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      CCCCCAGGTG TCACTGGACC CCATGAAGAA TGTCACCTGC GAGAATGGGC TGCCGGCCGT
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      GGTGAGTTGT GTGCCTGGGC AGGTCTTCAG CCCTGACGGA CCCTCGAGAT TCCGGAAAGC
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      ATACAAGCCA GAGCAACCCC TGGTGCGACT GAGAGGCGGT GCCTACATCG GGGAGGGCCG
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      GTCGGCCAGT GTGGTCTGCA GAGAGCTGGG CTTTGGGAGT GCCAAAGAGG CAGTCACTGG
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      CTCCCGACTG GGGCAAGGGA TCGGACCCAT CCACCTCAAC GAGATCCAGT GCACAGGCAA
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      GGATGCTGGT GTGAGATGCA ACACCCCTGC CATGGGCTTG CAGAAGAAGC TGCGCCTGAA
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      TGTGTGGGG ATGGTGTGT GCCAAAACTG GGGCATCGTG GAGGCCATGG TGGTCTGCCG
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      TGTCAACAGC AACAAGTGG TCATGAGTGG AGTGAAGTGC TCGGGAACGG AGCTGTCCCT
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      GGCGCACTGC CGCCACGACG GGGAGGACGT GGCCTGCCCC CAGGGCGGAG TGCAGTACGG
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      GGCCGGAGTT GCCTGCTCAG AAACCGCCCC TGACCTGGTC CTCAATGCGG AGATGGTGCA
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      GCAGACCACC TACCTGGAGG ACCGGCCCAT GTTCATGCTG CAGTGTGCCA TGGAGGAGAA
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      CTGCCTCTCG GCCTCAGCCG CGCAGACCGA CCCCACCACG GGCTACCGCC GGCTCCTGCG
      CTTCTCCTCC CAGATCCACA ACAATGGCCA GTCCGACTTC CGGCCCAAGA ACGGCCGCCA
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      CGCGTGGATC TGGCACGACT GTCACAGGCA CTACCACAGC ATGGAGGTGT TCACCCACTA
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      TGACCTGCTG AACCTCAATG GCACCAAGGT GGCAGAGGGC CACAAGGCCA GCTTCTGCTT
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      AAAGTTTGAG CACTTCAGCG GGCTCTTAAA CAACCAGCTG TCCCCGCAG<u>T AA</u>AGAAGCCT
      GCGTGGTCAA CTCCTGTCTT CAGGCCACAC CACATCTTCC ATGGGACTTC CCCCCAACAA
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      CTGAGTCTGA ACGAATGCCA CGTGCCCTCA CCCAGCCCGG CCCCCACCCT GTCCAGACCC
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      ACCTGACCCT TGGGGCCTGA GAAGGCCTTG GGGGGGTGGG GTTTGTCCAC AGAGCTGCTG
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      GAGCAGCACC AAGAGCCAGT CTTGACCGGG ATGAGGCCCA CAGACAGGTT GTCATCAGCT
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      TGTCCCATTC AAGCCACCGA GCTCACCACA GACACAGTGG AGCCGCGCTC TTCTCCAGTG
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      CTCCTCCTCT TAGGTCATTT TCAGCAAACT TGAATATCTA GACCTCTCTT CCAATGAAAC
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      GAAAGATTTA TG
      Seq ID No: 109 Protein sequence:
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                           21
                                      31
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                                                                          60
      AGOKRKHSEG RVEVYYDGOW GTVCDDDFSI HAAHVVCREL GYVEAKSWTA SSSYGKGEGP
                                                                         120
      IWLDNLHCTG NEATLAACTS NGWGVTDCKH TEDVGVVCSD KRIPGFKFDN SLINOIENLN
                                                                         180
      IQVEDIRIRA ILSTYRKRTP VMEGYVEVKE GKTWKQICDK HWTAKNSRVV CGMFGPPGER
                                                                         240
     TYNTKVYKMF ASRRKQRYWP FSMDCTGTEA HISSCKLGPQ VSLDPMKNVT CENGLPAVVS
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75
      CVPGQVFSPD GPSRPRKAYK PEQPLVRLRG GAYIGEGRVB VLKNGEWGTV CDDKWDLVSA
      SVVCRELGFG SAKEAVTGSR LGQGIGPIHL NEIQCTGNEK SIIDCKFNAE SQGCNHEEDA
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11

21

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GVRCHTPAMG LOKKLRENGG RNPYEGRVEV LVERNGSLVW GMVCGONWGI VEAMVVCRQL
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      GLGFASNAFO ETWYWHGDVN SNKVVMSGVK CSGTELSLAH CRHDGEDVAC POGGVOYGAG
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       VACSETAPDL VLNAEMVOOT TYLEDRPMFM LOCAMEENCL SASAAOTDPT TGYRRLLRFS
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       SQIHNNGQSD FRPKNGRHAW IWHDCHRHYH SMEVFTHYDL LNLNGTKVAE GHKASFCLED
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       TECEGDIQKN YECANFGDQG ITMGCWDMYR HDIDCQWVDI TDVPPGDYLF QVVINPNFEV
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      GCCCACTCTG GCTAGGGGGC GGCAGGCTGA GAGCTCACCT GTTCAGCAGA GAAGTGGAAC
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      CACTITECTC CTEGAGCCTG TCTACCACAG TGTTATCAGC TTCATTGTCA TCCTGGTGGT
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      GTGGTGATCA TCCTAGTTGG TGTGGTCAGC CTGAGGGTTC AGTGTCGGAA GAGCAAGGAG
                                                                           660
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      TCTGAAGATC CCAGAACCTG GGAGTACAGG GCGTGTCTGA CAAGCTGGTC ACAGACCATG
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      GGGC
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      Seq ID No: 111 Protein sequence:
      Protein Accession #: none found, CAT_73007_3
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      TSSSOGGLGG LSLTTEPVSS TODTSLPORL TGQAICPALV PRRRCPQQWK RRRHKQRHIS
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      TVPPNSTTMS LSMREDATIL PAPRORLCSL WLHLGWSRVE AHSG
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      Coding sequence: 37-3453 (underlined sequences correspond to start and stop codons)
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      ACGCTTCGCG GCTTCTCCAA GCCCTCGGAC CTCGTGGGCG TCTTCTCCTG CGTGGGCGGT
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      TCGAGCGGCA TCTACAGTGC CACTTACCTG GAAGCCAGCC CCCTGGGCAG CGCCTTCTTT
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      GACCCACTGT GGATAGACGT GGACAGGCCT GAGGAGACAA GCACCATCAT CCGTGGCCTC
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      AACGCCAGCA CGCGCTACCT CTTCCGCATG CGGGCCAGCA TTCAGGGGCT CGGGGACTGG
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      AGCTGCCTGC ATCGGAGACG CACCTTCACC TACCAGTCAG GCTCGGGCGA GGAGACCATC
      CTGCAGTTCA GCTCAGGGAC CTTGACACTT ACCCGGCGGC CAAAACTGCA GCCCGAGCCC
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      CTGAGCTACC CAGTGCTAGA GTGGGAGGAC ATCACCTTTG AGGACCTCAT CGGGGAGGGG
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      QLPNVQPPSS GIYSATYLEA SPLGSAFFRL IVRGCGAGRW GPGCTKECPG CLHGGVCHDH
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      DGECVCPPGF TGTRCEQACR EGRFGQSCQE QCPGISGCRG LTFCLPDPYG CSCGSGWRGS
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      EFNLETMPRI NCAAAGNPPP VRGSIELRKP DGTVLLSTKA IVEPEKTTAE FEVPRLVLAD
      SGFWECRVST SGGQDSRRPK VNVKVPPVPL AAPRLLTKQS RQLVVSPLVS FSGDGPISTV
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      RLHYRPQDST MDWSTIVVDP SENVTLMNLR PKTGYSVRVQ LSRPGEGGEG AWGPPTLMTT
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      DCPEPLLQPW LEGWHVEGTD RLRVSWSLPL VPGPLVGDGF LLRLWDGTRG QERRENVSSP
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      QARTALLTGL TPGTHYQLDV QLYHCTLLGP ASPPAHVLLP PSGPPAPRHL HAQALSDSEI
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      SIQGLGDWSN TVEESTLGNG LQAEGPVQES RAAEEGLDQQ LILAVVGSVS ATCLTILAAL
55
      LTLVCIRRSC LHRRRTFTYQ SGSGEETILQ FSSGTLTLTR RPKLQPEPLS YPVLEWEDIT
      FEDLIGEGNF GOVIRAMIKK DGLKMNAAIK MLKEYASEND HRDFAGELEV LCKLGHHPNI
      INLLGACKNR GYLYIAIEYA PYGNLLDFLR KSRVLETDPA FAREHGTAST LSSROLLRFA
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      SDAANGMQYL SEKQFIHRDL AARNVLVGEN LASKIADFGL SRGEEVYVKK TMGRLPVRWM
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      AIESLAYSVY TTKSDVWSPG VLLWEIVSLG GTPYCGMTCA ELYEKLPQGY RMEQPRNCDD 1080
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      EVYELMROCW RDRPYERPPF AOIALOLGRM LEARKAYVNM SLFENFTYAG IDATAEEA
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      Protein Accession #: NP_002623.1
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      DTKKCAFRTH TGKYWTLTAT GGVQSTASSK NASCYFDIEW RDRRITLRAS NGKFVTSKKN
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360

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⁷⁵ Seq ID NO: 126 <u>DNA sequence</u>
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Seq ID No: 135 Protein sequence:
Protein Accession #: XP_059648.1

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Seq ID NO: 136 DNA sequence

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Coding sequence: 304-2451 (underlined)

Coding sequence: 304-2451 (underlined sequences correspond to start and stop codons)

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      LGRDYSMVES PLICKEGESV QGSHVTRWPG FYILQWKFHS MPACAASSLP RVDDVLASLQ
                                                                           660
      VSSHKCKVMY YTEVIGSEDF RGSMTSLESS HSGFSQLSAA TTSSSQSHSS SMISR
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      Nucleic Acid Accession #: NM_004181.1
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      GGTCGCCGGC CAGTGGCGCT TCGTGGACGT GCTGGGGCTG GAAGAGGAGT CTCTGGGCTC
                                                                           120
      GGTGCCAGCG CCTGCCTGCG CGCTGCTGCT GCTGTTTCCC CTCACGGCCC AGCATGAGAA
                                                                           180
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      CTTCAGGAAA AAGCAGATTG AAGAGCTGAA GGGACAAGAA GTTAGTCCTA AAGTGTACTT
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      CATGAAGCAG ACCATTGGGA ATTCCTGTGG CACAATCGGA CTTATTCACG CAGTGGCCAA
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      TAATCAAGAC AAACTGGGAT TTGAGGATGG ATCAGTTCTG AAACAGTTTC TTTCTGAAAC
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      AGAGAAAATG TCCCCTGAAG ACAGAGCAAA ATGCTTTGAA AAGAATGAGG CCATACAGGC
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      AGCCCATGAT GCCGTGGCAC AGGAAGGCCA ATGTCGGGTA GATGACAAGG TGAATTTCCA
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      TTTTATTCTG TTTAACAACG TGGATGGCCA CCTCTATGAA CTTGATGGAC GAATGCCTTT
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      TCCGGTGAAC CATGGCGCCA GTTCAGAGGA CACCCTGCTG AAGGACGCTG CCAAGGTGTG
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      CAGAGAATTC ACCGAGCGTG AGCAAGGAGA AGTCCGCTTC TCTGCCGTGG CTCTCTGCAA
      GGCAGCC<u>TAA</u> TGCTCTGTGG GAGGGACTTT GCTGATTTCC CCTCTTCCCT TCAACATGAA
                                                                           720
                                                                           780
      AATATATACC CCCCATGCAG TCTAAAATGC TTCAGTACTT GTGAAACACA GCTGTTCTTC
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      TGTTCTGCAG ACACGCCTTC CCCTCAGCCA CACCCAGGCA CTTAAGCACA AGCAGAGTGC
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TATGTCTTGT ATCCGATATC TAACGCTTTA AATGGCTACT TTGGTTTCTG TCTGTAAGTT 960 AAGACCTTGG ATGTGGTTAT GTTGTCCTAA AGAATAAATT TTGCTGATAG TAGC

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      Protein Accession #: NP 004172.1
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      GQEVSPKVYF MKQTIGNSCG TIGLIHAVAN NQDKLGFEDG SVLKQFLSET EKMSPEDRAK
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      CFEKNEAIQA AHDAVAQEGO CRVDDKVNFH FILFNNVDGH LYELDGRMPF PVNHGASSED
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      TLLKDAAKVC REFTEREQGE VRFSAVALCK AA
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      GGCTCCGTGC TGGTGACATG CAGCACCTCC TGTGACCAGC CCAAGTTGTT GGGCATAGAG
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      ACCCCGTTGC CTAAAAAGGA GTTGCTCCTG CCTGGGAACA ACCGGAAGGT GTATGAACTG
                                                                          300
      AGCAATGTGC AAGAAGATAG CCAACCAATG TGCTATTCAA ACTGCCCTGA TGGGCAGTCA
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      CCCTCTTGGC AGCCAGTGGG CAAGAACCTT ACCCTACGCT GCCAGGTGGA GGGTGGGGCA
      CCCCGGGCCA ACCTCACCGT GGTGCTGCTC CGTGGGGAGA AGGAGCTGAA ACGGGAGCCA
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      GCTGTGGGGG AGCCCGCTGA GGTCACGACC ACGGTGCTGG TGAGGAGAGA TCACCATGGA
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      ATACAACTGG GAAATACTGA AACTTGCTGC CTATTGGGTA TGCTGAGGCC CACAGACTTA
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      CAGAAGAAGT GGCCCTCCAT AGACATGTGT AGCATCAAAA CACAAAGGCC CACACTTCCT
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      TACAGGTTGT ACACTGCAGG AGAGTGCCTG GCAAAAAGAT CAAATGGGGC TGGGACTTCT
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      CCTCCCAGCT TTGGAAGGGT CATCCGCGTG TGTGTGTGTG TGTATGTGTA GACAAGCTCT
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      TTTGGGCTCA AGTGATCCTC CCACCTCAGC CTCCTGAGTA GCTGGGACCA TAGGCTCACA
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      ACACCACAC TGGCAAATTT GATTITTTTT TTTTTTTCA GAGACGGGT CTCGCAACAT
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      Seq ID No: 141 Protein sequence:
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Protein Accession #: NP_000192.1

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      GTTCCACATG TATGAAGGGT ACCCACTCTG GAAGGTGACA TTCCCAGTGA GGGTTTTCCA
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      TGCCAGATCC TCTTCTCAAA GCTGGGATTA CAGGTGTGAG CATAGTGAGA CCTTGGCGCT
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      VPGHAGRLVF GFLNGRACVM MQGRFHMYEG YPLWKVTFPV RVFHLLGVDT LVVTNAAGGL
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      NPKFEVEDIM LIRDHINLPG PSGQNPLRGP NDERFGDRFP AMSDAYDRTM RQRALSTWKQ
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      MGEORELQEG TYVMVAGPSF ETVAECRVLQ KLGADAVGMS TVPEVIVARH CGLRVFGFSL
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      TTGAAAGCGA AGTTCAGGAA GAGTGACACC AATGAGTGGA ACAAGAATGA TGACCGGCTA
                                                                          180
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      CTGCAGGCCG TGGAGAATGG AGATGCGGAG AAGGTGGCCT CACTGCTCGG CAAGAAGGGG
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      GCCAGTGCCA CCAAACACGA CAGTGAGGGC AAGACCGCTT TCCATCTTGC TGCTGCAAAA
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      GGACACGTGG AATGCCTCAG GGTCATGATT ACACATGGTG TGGATGTGAC AGCCCAAGAT
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      ACTACOGGAC ACAGCGCCTT ACATCTCGCA GCCAAGAACA GCCACCATGA ATGCATCAGG
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      AGGCTGCTTC AGTCTAAATG CCCAGCCGAA AGTGTCGACA GCTCTGGGAA AACAGCTTTA
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      ATAAATGAGA TGTCGAAGGA AGTCACCAAA TTGAAGGAGG CCTTGAACAG CCTCTCCCAG
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                                                                          300
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      CACATGTGAG GAGCCTGAGT GCAACGACAT CACTGCCAGG CTGCAGTATG TCAAGGTGGG
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      ETEAGYYKLS GEAYGFVARI DGSGNFQVLL SDRYFNKTCG LCGNFNIFAE DDFMTQEGTL
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      TSDPYDFANS WALSSGEOWC ERASPPSSSC NISSGEMOKG LWEQCOLLKS TSVFARCHPL
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      VDPEPFVALC EKTLCECAGG LECACPALLE YARTCAQEGM VLYGWTDHSA CSPVCPAGME
      YROCVSPCAR TCQSLHINEM CQERCVDGCS CPEGOLLDEG LCVESTECPC VHSGKRYPPG
                                                                           360
      TSLSRDCNTC ICRNSOWICS NEECPGECLV TGQSHFKSFD NRYFTFSGIC QYLLARDCQD
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      HSPSIVIETV QCADDRDAVC TRSVTVRLPG LHNSLVKLKH GAGVAMDGOD IQLPLLKGDL
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      RICHTVTASV RLSYGEDLOM DWDGRGRLLV KLSPVYAGKT CGLCGNYNGN QGDDFLTPSG
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      LABPRVEDFG NAWKLHGDCQ DLQKQHSDPC ALNPRMTRFS EEACAVLTSP TFEACHRAVS
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      PLPYLRNCRY DVCSCSDGRE CLCGALASYA AACAGRGVRV AWREPGRCEL NCPKGQVYLQ
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      IFSDHHTMCY CEDGFMHCTM SGVPGSLLPD AVLSSPLSHR SKRSLSCRPP MVKLVCPADN
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      LRAEGLECTK TCQNYDLECM SMGCVSGCLC PPGMVRHENR CVALERCPCF HQGKEYAPGE
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      YIILLIGKAL SVVWDRHLSI SVVLKQTYQE KVCGLCGNFD GIQNNDLTSS NLQVEEDPVD
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      FGNSWKVSSQ CADTRKVPLD SSPATCHNNI MKQTMVDSSC RILTSDVFQD CNKLVDPEPY
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      LOVCIYOTCS CESIGDCACF COTIAAYAHV CAQHGKVVTW RTATLCPQSC EERNLRENGY
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      YHDGSHAYIG LKDRKRPSEL RRIASQVKYA GSQVASTSEV LKYTLFQIFS KIDRPEASRI
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      IDVPWNVVPB KAHLLSLVDV MQREGGPSQI GDALGFAVRY LTSEMHGARP GASKAVVILV
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      EQDLEVILHN GACSPGARQG CMKSIEVKHS ALSVELHSDM EVTVNGRLVS VPYVGGNMEV
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      NVYGAIMHEV RFNHLGHIFT FTPQNNEFQL QLSPKTFASK TYGLCGICDE NGANDFMLRD
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      DGNVSSCGDH PSEGCFCPPD KVMLEGSCVP EEACTQCIGE DGVQHQFLEA WVPDHQPCQI
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      WASPENPCLI NECVRVKEEV PIQQRNVSCP QLEVPVCPSG FQLSCKTSAC CPSCRCERME
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      ACMINGTVIG PGKTVMIDVC TTCRCMVQVG VISGFKLECR KTTCNPCPLG YKBENNTGEC
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                                                                           2700
      CGRCLPTACT IQLRGGQIMT LKRDETLQDG CDTHFCKVNE RGEYFWEKRV TGCPPFDEHK
      CLAEGGKIMK IPGTCCDTCB EPECNDITAR LQYVKVGSCK SEVEVDIHYC QGKCASKAMY
      SIDINDVQDQ CSCCSPTRTE PMQVALHCTN GSVVYHEVLN AMECKCSPRK CSK
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      ACGTCCAGCT ACACCCTGTC CTGCAAGCTG CACACTTTCC TCTTCGAGGC CTGCAGCTAC
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      CAGTCCAGCA TCTTCGGCGC CTTCGTGGTC TACCTCGTGG TCCTGCTCTC CGTAGCCTTC
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      ATTCGGAGGA TCATGGCTGC GGCCAAACCC AAGCACGACT GGACGAGGTC CTACTTCCGG
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      GCGTACATGA TCCTCCTCCC CTTCTCGGAG ACGTTTTTCT ACCTCAGCTC GGTCATCAAC
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      AATTCTGCTG CAGAGAATGG TTTTCAGGAG CATGAAGTTT GA
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70
      VNVPSHRGLT CNRSSTRHHE OPETSNMSIC TNLSSRWTVF QSSIFGAFVV YLVVLLSVAF
      MCWNMMQVLM KSQKGSLAGG TRPPQLRKSE SEESRTARRQ TIIFLRLIVV TLAVCWMPNQ
                                                                            300
      IRRIMAAAKP KHDWTRSYFR AYMILLPFSE TFFYLSSVIN PLLYTVSSQQ FRRVFVQVLC
                                                                            360
      CRLSLQHANH EKRLRVHAHS TTDSARFVQR PLLFASRRQS SARRTEKIFL STFQSEAEPO
      SKSOSLSLES LEPNSGAKPA NSAAENGFOR HEV
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Nucleic Acid Accession #: none found Coding sequence: 3-65 (underlined sequences correspond to start and stop codons)

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      CATGTAATTG AAATTCATGC AAGGAAACAA CTCATAGAAA AGATAAATAT GGATGCCCTT
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                                                                        300
      TAAATGACAA AACACAGCAG TGTTTTGAGG GAGAAAGGAC CATCATTTAT AATGCTCTGT
                                                                        360
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      TTAAGCCCTT TTGCATTTGC CAGCCGTTGA CATTAAGAGG CATGTTTAAC GGTGCCAACA
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	CGCACTTTTA	TTTGTATTTT	TTCAGATTTT	TTTTTGTTTC	GTGGTGGTGG	GGGAGGTGAT	780
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		TTGCTCTGGA					900
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	GAGCGCCACC	GACCGGGATG	AGGGCGCCAA	CGCCCAGCTT	GCCTACTCTA	TCCTCGAGTG	2340
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		TCGATAAATT					4980

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      TRVAAVDADD GENARLTYSI VRGNEMNLFR MDWRTGELRT ARRVPAKRDP QRPYELVIEV
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      Coding sequence: 70-1890 (underlined sequences correspond to start and stop codons)
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      AATGATGGAA CCGTCTGTAT AGAAAATGTG AATGCAAACT GCCATTTAGA TAATGTCTGT
                                                                          180
      ATAGCTGCAA ATATTAATAA AACTTTAACA AAAATCAGAT CCATAAAAGA ACCTGTGGCT
                                                                          240
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      TTGCTACAAG AAGTCTATAG AAATTCTGTG ACAGATCTTT CACCAACAGA TATAATTACA
      TATATAGAAA TATTAGCTGA ATCATCTTCA TTACTAGGTT ACAAGAACAA CACTATCTCA
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      GCCAAGGACA CCCTTTCTAA CTCAACTCTT ACTGAATTTG TAAAAACCGT GAATAATTTT
                                                                          420
      GTTCAAAGGG ATACATTTGT AGTTTGGGAC AAGTTATCTG TGAATCATAG GAGAACACAT
                                                                          480
      CTTACAAAAC TCATGCACAC TGTTGAACAA GCTACTTTAA GGATATCCCA GAGCTTCCAA
                                                                          540
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      AAGACCACAG AGTTTGATAC AAATTCAACG GATATAGCTC TCAAAGTTTT CTTTTTTGAT
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      TCATATAACA TGAAACATAT TCATCCTCAT ATGAATATGG ATGGAGACTA CATAAATATA
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      TTTCCAAAGA GAAAAGCTGC ATATGATTCA AATGGCAATG TTGCAGTTGC ATTTTTATAT
                                                                          720
      TATAAGAGTA TIGGTCCTTT GCTTTCATCA TCTGACAACT TCTTATIGAA ACCTCAAAAT
                                                                          780
      TATGATAATT CTGAAGAGGA GGAAAGAGTC ATATCTTCAG TAATTTCAGT CTCAATGAGC
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      TCAAACCCAC CCACATTATA TGAACTTGAA AAAATAACAT TTACATTAAG TCATCGAAAG
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      GTCACAGATA GGTATAGGAG TCTATGTGCA TTTTGGAATT ACTCACCTGA TACCATGAAT
                                                                          960
      GGCAGCTGGT CTTCAGAGGG CTGTGAGCTG ACATACTCAA ATGAGACCCA CACCTCATGC
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      CGCTGTAATC ACCTGACACA TTTTGCAATT TTGATGTCCT CTGGTCCTTC CATTGGTATT
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      AAAGATTATA ATATTCTTAC AAGGATCACT CAACTAGGAA TAATTATTTC ACTGATTTGT
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      CTTGCCATAT GCATTTTTAC CTTCTGGTTC TTCAGTGAAA TTCAAAGCAC CAGGACAACA
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      ATTCACAAAA ATCTTTGCTG TAGCCTATTT CTTGCTGAAC TTGTTTTTCT TGTTGGGATC
      AATACAAATA CTAATAAGCT CTTCTGTTCA ATCATTGCCG GACTGCTACA CTACTTCTTT
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      TTAGCTGCTT TTGCATGGAT GTGCATTGAA GGCATACATC TCTATCTCAT TGTTGTGGGT
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      GTCATCTACA ACAAGGGATT TTTGCACAAG AATTTTTATA TCTTTGGCTA TCTAAGCCCA
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      ATTCTTGTTA ATCTCTTGGC TTTTGGAGTC ATCATATACA AAGTTTTTCG TCACACTGCA
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      GGGTTGAAAC CAGAAGTTAG TTGCTTTGAG AACATAAGGT CTTGTGCAAG AGGAGCCCTC
      GCTCTTCTGT TCCTTCTCGG CACCACCTGG ATCTTTGGGG TTCTCCATGT TGTGCACGCA
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      TCAGTGGTTA CAGCTTACCT CTTCACAGTC AGCAATGCTT TCCAGGGGAT GTTCATTTTT
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      TTATTCCTGT GTGTTTTATC TAGAAAGATT CAAGAAGAAT ATTACAGATT GTTCAAAAAT
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                                                                        1920
      ACAAAATAA AAATTCCAAG CTGTGGATGA CCAATGTATA AAAATGACTC ATCAAATTAT
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      AACTGTAGAT AATAAGGTAA AATTATGTAT CATATAGATA TACTATGTTT TTCTATGTGA
      AATAGTTCTG TCAAAAATAG TATTGCAGAT ATTTGGAAAG TAATTGGTTT CTCAGGAGTG
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      CAAACAGTGA AAAGGGAATG ATAAGATGTA TTTTGAATGA ACTGTTTTTT CTGTAGACTA
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      GCTGAGAAAT TGTTGACATA AAATAAAGAA TTGAAGAAAC ACATTTTACC ATTTTGTGAA
      TTGTTCTGAA CTTAAATGTC CACTAAAACA ACTTAGACTT CTGTTTGCTA AATCTGTTTC
      TTTTTCTAAT ATTCTAAAA
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      Protein Accession #: NP 071442.1
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      EVYRNSVTDL SPTDIITYIE ILAESSSLLG YKNNTISAKD TLSNSTLTEF VKTVNNFVQR
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      DTFVVWDKLS VNHRRTHLTK LMHTVEQATL RISQSFQKTT BFDTNSTDIA LKVFFFDSYN
                                                                             180
      MKHIHPHMNM DGDYINIFPK RKAAYDSNGN VAVAFLYYKS IGPLLSSSDN FLLKPONYDN
                                                                             240
      SEEEERVISS VISVSMSSNP PTLYBLEKIT FTLSHRKVTD RYRSLCAFWN YSPDTMNGSW
20
      SSEGCELTYS NETHTSCRCN HLTHFAILMS SGPSIGIKDY NILTRITQLG IIISLICLAI
                                                                             360
      CIFTFWFFSE IQSTRTTIHK NLCCSLFLAE LVFLVGINTN TNKLFCSIIA GLLHYFFLAA
                                                                             420
      FAWMCIEGIH LYLIVVGVIY NKGFLHKNFY IFGYLSPAVV VGFSAALGYR YYGTTKVCWL
                                                                             480
      STENNFIWSF IGPACLIILV NLLAPGVIIY KVFRHTAGLK PEVSCFENIR SCARGALALL
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      FLLGTTWIFG VLHVVHASVV TAYLFTVSNA FQGMFIFLFL CVLSRKIQEE YYRLFKNVPC
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      Nucleic Acid Accession #: none found
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      GTATCTTCAA AATCCAGAAG GATGATGGCA GATGGCAGGA AGGAGGAAGA GGGTAATCTG
                                                                             180
      GAAGAGTTTC CTGACCTACT CTGCTGCTGT GAT<u>TAA</u>ACAA CCACCAGGAA ATTTTGATGA
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      CACTGTTCTC CTGAGCTCCT CCCTTTCCTC GGGGAAGAAA AGCATTGAAA CTACAAAAAT
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      AAAGTGTTAT TTGGCTGGAG TGAGGTCTCA TGTCTGCTTA TGCGGTGGCT CGCTGCTCAG
      AACAGGGAAC CATTGGAGAT ACTCATTACT CTTTGAAGGC TTACAGTGGA ATGAATTCAA
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      ATACGACTTA TTTGAGGAAT TGAAGTTGAC TTTATGGAGC TGATAAGAAT CTTCTTGGAG
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      AAAAAAAGAC TGGTACTTCT GAATTAACCA AAATCACAGT ATTCTGAAGA TGATTCTACA
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      AAGCCTGCTG TTTCTACAAA GGCTGCTGAT GATTTCTACA AAGCCTGCTG TAGTGTTGCT
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      GTGGCCTCTG CTTAAAAAAG TAGAAAACAC ATTGATGCAG CATGTTCACC CCAACCTCCC
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      TGCCTAAAGG CTCAGGGACC ATCTTGGAAG AGGAAGGCGC GTGAGATTGT AAGAGCCGAA
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      TTAGGGGGAT GGAGTGTGGA GAATAAGGAC ACTTCATCTT GGATGCTCAC CTGCCAAATT
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      GACTTCTGAT GAAAGCCAGC TCCAGAAATG TGCCTACAGT TACTACTTTC ACCTAAACCC
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      TGCCCTTAGT CAAATCCTTC TCTTCTTCTA AGCAATCAAC TTCAATTCCT TGTATAACCC
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      ACAGTATAAA AGGGCTTTTA TACCATTCTA TCCTATTGCA TGTAAGCCTT GGGTCTGGGA
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      GGTAACAGTG TGGGATTCCA CCATCTCATC TCCCTGCCAC CCAAACATGC CTGCTCTTCT
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      TTAAGCAATA TTAAATGTTT GTACTTCA
      Seq ID No: 161 Protein sequence:
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      Protein Accession #: none found
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      CLLMRWLAAQ NRMAEMSTTI KNSRTSAVGP VICFMEFTSL VSSKSRRMMA DGRKEEBGNL
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      EEFPDLLCCC D
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      Nucleic Acid Accession #: none found
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      Coding sequence: 1-159 (underlined sequences correspond to start and stop codons)
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      TCAGGAAGGA CCCACAGACT GCTGCTCCTG GAGGCCTCTC GGTTTATGGA TGTGTGTTTG
                                                                             120
      TTCCATAAAC CCTCAGAGGG TCACCTGGAG ACCCGCTAAA ATGCAGGTTC TTGGGCCACA
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      TCCTAGACCT TCTGACCGAC CCAGGGAGTG GGGCCCAGGA AGCTGCATTT GACAGATATC
                                                                             240
      CCCGTGTGAT CATCATGCAC ACAGGAGTGA GAGAACCAGT GTTCTCCCCG GGCAGAAGGG
                                                                             300
75
      AAGCTCGTGT GCAGGACACC TCACACCTCC TTTCCCATTC CCCTGCCAGG CTCTCCCTGC
                                                                             360
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TGACATTGTT TTTGCGGGAG AGCTGTGAAT TCTGAAGATT AGGTTGCTTC TCACCCCAAG

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CTCCAGAAGT CCAGGCTGAG CCAAACCAAG CTTCAAGTTG TGCCTGGACT TGGAGAACCA
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                                                                           540
      GCACCAAGGA AAGGCCATGC AAGGACACAG GGAGAAGGGC AGCTGTCTGT AAGCCAGAAA
      GAGCCTTCAC TAGAAACCAA ATCAGCCAGA ACCTTCATCT TGGACTTTCC AGCCTTCAGA
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      GATGTGAAAA AATAAATTTC TGTTGATTAA CCTAAAAAA
      Seq ID No: 163 Protein sequence:
      Protein Accession #: none found
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      ETLOROGPGL KREALLHTCS SGRTHRLLLL EASRFMDVCL PHKPSEGHLE TR
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      GACTACCTGA ACCACTATCC CGTGTTTGTG GGCAGCGGGC CCGGACGCCT GACCCCCGCA
25
      GAAGGTGCTG ACGACCTCAA CATCCAGCGA GTCCTGCGGG TCAACAGGAC GCTGTTCATT
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      GGGGACAGGG ACAACCTCTA CCGCGTAGAG TTGGAGCCCC CCACGTCCAC GGAGCTGCGG
                                                                           300
      TACCAGAGGA AGCTGACCTG GAGATCTAAC CCCAGCGACA TAAACGTGTG TCGGATGAAG
GGCAAACAGG AGGGCGAGTG TCGAAACTTC GTAAAGGTGC TGCTCCTTCG GGACGAGTCC
                                                                           360
                                                                           420
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      ACCCTGCAGC CCGTCGGAGA CAACATCAGC GGTATGGCCC GCTGCCCGTA CGACCCCAAG
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      CTAGCCATTG ATGCTGTCAT CTACCGCAGC CTCGGGGACA GGCCCACCCT GCGCACCGTG
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      AAACATGACT CCAAGTGGTT CAAAGAGCCT TACTTTGTCC ATGCGGTGGA GTGGGGCAGC
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      CATGTCTACT TCTTCTCCG GGAGATTGCG ATGGAGTTTA ACTACCTGGA GAAGGTGGTG
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      GTGTCCCGCG TGGCCCGAGT GTGCAAGAAC GACGTGGGAG GCTCCCCCCG CGTGCTGGAG
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      TTCTACTTCA ACGTGCTGCA GGCTGTCACG GGCGTGGTCA GCCTCGGGGG CCGGCCCGTG
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      GTCCTGGCCG TTTTTTCCAC GCCCAGCAAC AGCATCCCTG GCTCGGCTGT CTGCGCCTTT
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      1140
      GCCCCCGGGA TGCAGTACAA TGCCTCCAGC GCCTTGCCGG ATGACATCCT CAACTTTGTC
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      AAGACCCACC CTCTGATGGA CGAAGCGGTG CCCTCGCTGG GCCATGCGCC CTGGATCCTG
                                                                         1260
      CGGACCCTGA TGAGGCACCA GCTGACTCGA GTGGCTGTGG ACGTGGGAGC CGGCCCCTGG
                                                                         1320
      GGCAACCAGA CCGTTGTCTT CCTGGGTTCT GAGGCGGGGA CGGTCCTCAA GTTCCTCGTC
                                                                          1380
45
      CGGCCCAATG CCAGCACCTC AGGGACGTCT GGGCGTGTGT GTCAAGTGGG CCACGCGTGC
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      AGGGTGTGTG TCCACGAGCG ACGATCGTGG TGGCCCCAGC GGCCTGGGCG TTGGCTGAGC
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                                                                         1560
      CCCCCACTC TGCAGAGGGA AGCGGGGACA ATGCCGGGGT TTCAGGCAGG AGACACGAGG
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      AGGGCCTGCC CGGAAGTCAC ATCGGCAGCA GCTGTCTAAA GGGCTTGGGG GCCTGGGGGG
                                                                         1680
50
      CGGCGAAGGT GGGTGGGGCC CCTCTGTAAA TACGGCCCCA GGGTGGTGAG AGAGTCCCAT
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                                                                           60
60
      GADDLNIQRV LRVNRTLFIG DRDNLYRVEL EPPTSTELRY QRKLTWRSNP SDINVCRMKG
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      KQEGECRNFV KVLLLRDEST LFVCGSNAFN PVCANYSIDT LQPVGDNISG MARCPYDPKH
      ANVALFSDGM LFTATVTDFL AIDAVIYRSL GDRPTLRTVK HDSKWFKEPY FVHAVEWGSH
                                                                          240
      VYFFFREIAM EFNYLEKVVV SRVARVCKND VGGSPRVLEK QWTSFLKARL NCSVPGDSHF
                                                                          300
      YFNVLQAVTG VVSLGGRPVV LAVFSTPSNS IPGSAVCAFD LTQVAAVFEG RFREOKSPES
                                                                          360
65
      IWTPVPEDQV PRPRPGCCAA PGMQYNASSA LPDDILNFVK THPLMDEAVP SLGHAPWILR
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      TLMRHQLTRV AVDVGAGPWG NQTVVFLGSE AGTVLKFLVR PNASTSGTSG RVCQVGHACR
                                                                          480
      VCVHERRSWW PQRPGRWLSR RWGFQKARGP PRCRLGV
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180

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      ACTTOGTAAA GGTGCTGCTC CTTCGGGACG AGTCCACGCT CTTTGTGTGC GGTTCCAACG
                                                                           480
      CCTTCAACCC GGTGTGCGCC AACTACAGCA TAGACACCCT GCAGCCCGTC GGAGACAACA
      TCAGCGGTAT GGCCCGCTGC CCGTACGACC CCAAGCACGC CAATGTTGCC CTCTTCTCTG
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                                                                          840
      AGAACGACGT GGGAGGCTCC CCCCGCGTGC TGGAGAAGCA GTGGACGTCC TTCCTGAAGG
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      GTTCTGAGGC GGGGACGGTC CTCAAGTTCC TCGTCCGGCC CAATGCCAGC ACCTCAGGGA
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                                                                         1680
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                                                                         1980
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      GCTGGGCCAA GGCCACGCTG CTGCAGGGCG GGCCCCACGA CCTGGACTCG GGGCTGCTGC
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      ACGCCCTGGG CCCCCGCGCC TGGGACCACG GCCACCCCCT GCTCCCGGCC TCCGCTTCAT
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      CCTCCCTCT GCTGCTGGCG CCCGCCGGG CCCCCGAGCA GCCCCCGCG CCTGGGGAGC
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      CCTAGGCCGG GGGCCCCCCG ATGCCTTGGC AGTGCCAGCC ACGGGAACCA GGAGCGAGAG
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      CTTTTGCGGT TTTCTAACCA ATTGCACAAC TCCGTTCTCG GGGTGGCGGC AGGCAGGGGA
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      Protein Accession #: NP_115484.1
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                                                                          120
      KQBGECRNPV KVLLLRDEST LFVCGSNAFN PVCANYSIDT LQPVGDNISG MARCPYDPKH
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                                                                          240
      VYFFFREIAM EFNYLEKVVV SRVARVCKND VGGSPRVLEK QWTSFLKARL NCSVPGDSHF
                                                                          300
      YFNVLQAVTG VVSLGGRPVV LAVFSTPSNS IPGSAVCAFD LTQVAAVFEG RFREQKSPES
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      TLMRHQLTRV AVDVGAGPWG NOTVVFLGSE AGTVLKFLVR PNASTSGTSG LSVFLREFET
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      YRPDRCGRPG GGETGORLLS LELDAASGGL LAAFPRCVVR VPVARCQQYS GCMKNCIGSQ
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      TSSVAAFVVG AVVSGFSVGW FVGLRERREL ARRKDKEAIL AHGAGEAVLS VSRLGERRAQ
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      GTCTGCTGGA CCCCAGCACA GGTCTGGCCC TGGACCGGCT GAGCGCCCCT GACCCGGCCT
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      GGATGGCGAG ACTCTCTTG CCCCTCACCA CCAACTACCG TGACAATGTG ATCTCCCCGG
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      GGCGGCTCTC GGTCTGCGGG AGGACCCTCA GTTTAGACTT GGCCACCAGT GCAGCCTCAG
      GCATGAAAGT GCAAGGGGAC CCAGGTGGAA AGACGGGGAC TGAGGGCAAG AGCAGAGGCA
      GCAGCAGCAG CAGCAGGTGC CTGTGAACAT ACCTCAGACG CCTCTGGATC CAAGAACCAG
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      GGGCCTGAGG ATCTGTGGAC AAGAGCTGGT TTCTAAAATC TTGTAACTCA CTAGCTAGCG
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      GCGGCCTGAG AACTTTAGGG TGACTGATGC TACCCCCACA GAGGAGGCAA GAGCCCCAGG
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                                                                         4980
      ACTAACAGCT GACTGACCAA AGCAGCCCCT TGTAAGCAGC TCTGAGTCTT TTGGAGGACA
      GGGACGGTTT GTGGCTGAGA TAAGTGTTTC CTGGCAAAAC ATATGTGGAG CACAAAGGGT
      CAGTCCTCTG GCAGAACAGA TGCCACGGAG TATCACAGGC AGGAAAGGGT GGCCTTCTTG
      GGTAGCAGGA GTCAGGGGGC TGTACCCTGG GGGTGCCAGG AAATGCTCTC TGACCTATCA 5160
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      ATAAAGGAAA AGCAGTGATT CAAAAAAAAA AAAAAAAAA AAAAAAAAA
      Seq ID No: 175 Protein sequence:
      Protein Accession #: NP_057664.1
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                            21
                                       31
                                                  41
                                                             51
      MMOLLOLLIG LIGPGGYLFL LGDCOEVITL TVKYQVSEEV PSGTVIGKLS QELGREERRR
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      QAGAAPQVLQ LPQALPIQVD SEEGLLSTGR RLDREQLCRQ WDPCLVSFDV LATGDLALIH
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      VEIQVLDIND HQPRFPKGEQ ELEISESASL RTRIPLDRAL DPDTGPNTLH TYTLSPSEHF
                                                                          180
      ALDVIVGPDR TKHABLIVVK ELDREIHSFF DLVLTAYDNG NPPKSGTSLV KVNVLDSNDN
      SPAFAESSLA LEIGEDAAPG TLLIKLTATD PDQGPNGEVE FFLSKHMPPE VLDTFSIDAK
      TGQVILRRPL DYEKNPAYEV DVQARDLGPN PIPAHCKVLI KVLDVNDNIP SIHVTWASQP
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      SLVSEALPKD SFIALVMADD LDSGHNGLVH CWLSOELGHF RLKRTNGNTY MILITNATIOR
                                                                          420
65
      EQWPKYTLTL LAQDQGLQPL SAKKQLSIQI SDINDNAPVF EKSRYEVSTR ENNLPSLHLI
                                                                          480
      TIKAHDADLG INGKVSYRIQ DSPVAHLVAI DSNTGEVTAQ RSLNYEEMAG FEFQVIAEDS
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      GQPMLASSVS VWVSLLDAND NAPEVVQPVL SDGKASLSVL VNASTGHLLV PIETPNGLGP
                                                                          600
      AGTDTPPLAT HSSRPFLLTT IVARDADSGA NGEPLYSIRS GNEAHLFILN PHTGQLFVNV
      TNASSLIGSE WELEIVVEDQ GSPPLQTRAL LRVMFVTSVD HLRDSARKPG ALSMSMLTVI
70
      CLAVLIGIFG LILALFMSIC RTEKKDNRAY NCREAESTYR QQPKRPQKHI QKADIHLVPV
      LRGQAGEPCE VGQSHKDVDK EAMMEAGWDP CLQAPFHLTP TLYRTLRNQG NQGAPAESRE
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      VLQDTVNLLF NHPRQRNASR ENLNLPEPQP ATGQPRSRPL KVAGSPTGRL AGDQGSERAP
                                                                          900
      QRPPASSATL RRQRHLNGKV SPEKESGPRQ ILRSLVRLSV AAFAERNPVE BLTVDSPPVQ
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      QISQLLSLLH QGQPQPKPNH RGNKYLAKPG GSRSAIPDTD GPSARAGGQT DPEQEEGPLD
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75
      PEEDLSVKQL LEEELSSLLD PSTGLALDRL SAPDPAWMAR LSLPLTTNYR DNVISPDAAA
      TEEPRTFQTF GKARAPELSP TGTRLASTFV SEMSSLLEML LEQRSSMPVE AASKALRRLS
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VCGRTLSLDL ATSAASGMKV QGDPGGKTGT EGKSRGSSSS SRCL

Seq ID NO: 176 DNA sequence

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Nucleic Acid Accession #: AL109712.1
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      Coding sequence: 2-128 (underlined sequences correspond to start and stop codons)
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                                       31
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                                                                           120
      CTGTGTGAAC TCTGGCAGGC TGCCCTCTCT GTGCAAAGCT GCCACTGGGG CCTGCTCAGG
      GTGGCCTGGA ACTTGGAGGT GGGCAGTCAG GGCCTAGGAT GGGCCTGTGT CACCAGGGCA
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      TGTGCCCTTG GGCCAGTTAC TTCCTCTCAG AGCCTTGGGC TCCTCCTCTG AGGATGGGGC
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      GCACAGCGGG CCAGCCTGCT CTCTTCTCTG TCTACTTTTT GCAGAAGAGT CAACAGATAC
      AACAGGCCCA GGGAGGTGCC CCTGGGGGCC CCAGTCCCCA TCACTCCAAG GGGCAGTCCT
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      GCAAGTGACA AGGTGGGCCC AATCCCTGTG GAACAGGTCT CTGAGGACCA CAGAGTGGGG
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      CCTGCCCGGA AGACCTTCCT GCCCCCAGAA CCCGACCCTC CGCAGATAGC CCTCCCTGGG
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      CAGCAGCCCC CCAGCTTCCA AGGCCCGTGC CTCACCAGAC GCCATGCTCT CACGGACTTG
                                                                           780
      TTTGCTGCTC TGTACCCTGC AGATCTGCCC CAGAGGAGCA GGTGAAAAGC CGCGCCTGCC
                                                                           840
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                                                                           900
25
      AGATICTCCA AATCCAAGAT GAAGTCATGC TGTGCTTTGG AATGGTAGAT GCTCATTTAT
      GTAAAATCAT AATAAATGTT ACACAAACTG TTAAAAAAAA AAAAAAAAA AAAAAA
      Seq ID No: 177 Protein sequence:
      Protein Accession #: AL109712.1
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                                                              51
                                                  41
      VSLGOPGCCR OTGSTPDAPL PSGSTAGLGL TLACPATCFP V
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      Coding sequence: 3-107(underlined sequences correspond to start and stop codons)
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      AAAGCATGCA ACTCTCCAGG GAACCAGAAC TAAATTGCTT AAAATGAAGT CATTCCTCAG
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      ATTACTTCC TCAGATAAAG TGTCAGCGGT CTGCAGAAAC GAAGAAGACA AAACTGAGAT
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      TATCACTCAT AATTCTCTTA CTTACTATGT CAGTGAAACA ATGAGTTTGC ATTTTTGCAA
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      CCTTTAGGAA AGATTTAAGA TGAACCTTCA ATAGGATATT AACATAACTC ATAGCCAATA
      CCACAGCTGC CTTTCAAATT AATGAGGTTA ATTGTTCTCC AGCAAACATG AGTTTGTCTT
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      GCCATCTAAG ACTTATTTAA TTATTTCTGT TCTCAGTCAA GCTAATTCAA GTGAATGAAC
                                                                           660
      AGTATTGACT TITAAAATCI TITITAAATT TITITAAATC TITAGTITAT TAAGTITGTA
                                                                           720
      GAAAAGCTCT GGGGCCATGA CCACTTACGT AAATGTTTCA GTTTAAAAAC AAAAGATTCA
                                                                           780
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      GGCCTCTAAT TTGAGCCAAA TCCAGGTGAT CTTGTTTGAA ATTTTTGATG AATTTGAAAA
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      ATTGCCTACT TAAGATATAA GTTCAAGAAT AACATTTTCA TAGAAAATTC AGAAAACTGC
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      TTGACACAGC AGTGACATAG TTAGATGTGG CTCAGATGCC TTCCAAACCT GAGGGTCCCC
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      AAAGATTTCT TTACCAGTTG TTTTTAACTA TGAATCTTAA TCTTGTTCAT TCCCCTGCCA
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      Seq ID No: 179 Protein sequence:
      Protein Accession #: none found
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COGGTGGGG CCTCGGGATG CAGGCGCCGG TGCCCGGGCC CCTGGGCCTG CTGGACCCCG
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      TGGGGGCTAC TACCCAGGGA TCATTCTCGG CTTTGGATCT TTCTTAGGAA TTATTGGCAT
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      CAACTTGGTG GAGAATAGAA GGCAAATGCT GGTGGCAGCG ATCGTGTTTA TCAGTTTTGG
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      CGATGTCTAC CAGACAGAGG TGAGCAGGAG CACTGAGATT CATGTGGGTT TTGCTCAGCT
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      TAATAGTTCC TGGGGGTCAC CCCTGCAGCT GGTGAACCGT TGATGCCCCC TGTGTAAGGG
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      Protein Accession #: none found
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      RVGPRDAGAG ARAPGPAGPR RRAFEEEEDV ALVCGVSAAG VRPHSHRRAG CISRTENV
      Seq ID NO: 182 DNA sequence
      Nucleic Acid Accession #: AK001579.1
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      GCCACCAGCT GCTGGGCCCC GGGCTGCTGC GGCTGGGCCG CCTATGGCTG CGGTCCCCCT
                                                                          180
      CCCATACAGC CCCGGCCCCT GGTCTCTGGC TGTCAGGGTT TGGCCTCCTT CGTGGTGACC
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      AGCTGCCACT CTTTCTTCCC GGGTCCTCCC CAACCCTCCT CCATTCCATC CCCAGAGCTG
                                                                         1020
                                                                         1080
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      CTAAACCAGA TGTGCACGCG GAACTTGGCT CTGCTGTTTG CACCCAGCGT GTTCCAGACG
                                                                         1200
      GATGGGCGAG GGGAGCACGA GGTGCGAGTG CTGCAAGAGC TCATTGATGG CTACATCTCT
                                                                         1260
      GTCTTTGATA TCGATTCTGA CCAGGTAGCT CAGATTGACT TGGAGGTCAG TCTTATCACC
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                                                                         1320
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      AGGITCTITC TGCTGCGTGG CCGCTGCCTG CTGCTGCTCA AGGAGAAGAA AAGCTCTAAA
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      GACCAGCAGC CAGTGGTCTT ACGACGCCAT TCCTCCTCTG ACCTTGCCCG TCAGAAGTTT
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      AATCAGACCC TGCGGCGACT ACACAACCGG AGGACCCTGT CCATGTTCTT TCCAATGAAG
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                                                                         2340
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                                                                         2400
      CCAGGCCCC CTTCAAAGAG CAGTCCCCAG GCACGGGGGT CCCTAGAGGA ACAGCTGCTC
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                                                                         2820
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      Seq ID No: 183 Protein sequence:
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      NCVTLKVSPT LTAEELTNQV LEMRGTAAGM DLWVTFEIRE HGELERPLHP KEKVLEQALQ
      WCQLPEPCSA SLLLKKVPLA QAGCLFTGIR RESPRVGLLR CREEPPRLLG SRFQERFFLL
                                                                          360
      RGRCLLLLKE KKSSKPEREW PLEGAKVYLG IRKKLKPPTP WGFTLILEKM HLYLSCTDED
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      EMWDWTTSIL KAQHDDQQPV VLRRHSSSDL ARQKFGTMPL LPIRGDDSGA TLLSANQTLR
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      RLHNRRTLSM FFPMKSSOGS VEEOEBLEEP VYEEPVYEEV GAFPELIODT STSFSTTREW
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      TVKPENPLTS QKSLDQPFLS KSSTLGQEER PPEPPPGPPS KSSPQARGSL EEQLLQELSS
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      GCACTTAAAA TAGAAAAAAA AAAAAAAAA AGCTCAGAGA GGCTGAGTTG TCCAAGGTCA
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_	CTATCCACTG	CCACATGTGA	AAAACTAAAT	TTGCAAACTT	CAACAAGCAT	TCCAAATGTT	360
5	ACTGAGATGA	AAACCAACAT	GAGCCTCGGC	TTGATCCTCA	CCAGGAACAT	GGGAACTGGA	420
-				CAGCAATGTG			480
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10				AAATTTGTAC			660
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	CARCCACCTC	<b>ምስርምስር</b> ስ አስከ	CCCTCCCACA	TTAGGAGAAC	A A A THEFT OF A	CATTICALGOT	1140
	ACIGITCAAG	GAGGAGACAA	CTTTCAGATG	GAAATGTCAC	AAGTGGGATT	CAGTGCAGAT	1200
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20							
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							1500
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				TTGGGTCAGC			1680
	GAGGGCATTG	AAAACACTCG	ATTTGGTTCA	GCAATTGCAG	CTCTTTCAGA	CATCAACATG	1740
	CATCCCTTTA	יי עניאבאי עניאי עניאי	TATALATICAL TATALATICAL TO A STATE OF THE ST	CCACTAGAAA	ልጥሮአርአልጥጥሮ	TOCACOTOTA	1800
	TACATTTACA	ATGGTCATCA	GGGCACTATC	CGCACAAAGT	ATTCCCAGAA	AATCTTGGGA	1860
30	TCCGATGGAG	CCTTTAGGAG	CCATCTCCAG	TACTTTGGGA	GGTCCTTGGA	TGGCTATGGA	1920
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	TCAACGTTCC	AGACAGTACA	GCTAACGGCA	GCTGCAGAAA	TCAACACCTA	TAACCCTGAG	3360
55	מבערבערמידער	TTGAAGATAA	בא ע האליאה ארני	PALALCIC CALCE P	ТСАТАВТОВ	ACCTGATGAG	3420
						CCTTTTGCTG	
	TTAGCTCTGG	TTGCAATTTT	ATGGAAGCTC	GGCTTCTTCA	AAAGAAAATA	TGAAAAGATG	3540
	ስርርል <u>ልልልል</u> ቸር	CACATCACAT	TOTATION CACC	ACAGAGCTCA	CTACCTCAAC	CAGCAGACCT	3600
<b>60</b>				CAGGGTTTGC			3660
60	TTTTTAAATC	CCATATTTTT	TTTATCATGT	CGTAGGTAAA	CTAACCTGGT	ATTTTAAGAG	3720
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				TATATGATGG			3840
				TTGCATTGTG			3900
	CAAGCATGAC	מ מ מידידידים א	CARARATOR	ATACTCTCAG	ATTITIANCCC	GGAAAACTGT	3960
65							
UJ						TGATATGTAA	
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	VCVCCtandalan.	ביד בידי בידי בידי	CTGCTCATCC	AAAGTTGCCA	CAGATGATAC	TTCCAAGTGA	4140
				TTGTTGGTTC			4200
	CTTCCACACC	CCATCTTGCT	CTAATGATCA	AAACATGCTT	GAATAACTGA	GCTTAGAGTA	4260
70	עס להולהלהלהוש ער	משופה עה ארולה לאות ע	PPCLLDGGDG	AGGGGGGGGAAT	<b>אדאכאכאריי</b> א	AGGCACAAAA	4320
				TGTAAAATCC			4380
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,				AACCTAGTGA			4500
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	TATTATAGAA	GCCCTCTACA	GCCTGACTTT	CTCTCCAGCG	GTCCAAAGTT	ATCCCCTCCT	4620
						GATATTAGGG	
							-000

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      GGCCGAGGCG AGCCGCGCAA GTTTCCCGGG ACCCTCAGAG TTGCACTCCG AAGACTCCAG
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      ATTCCGAGAG TTGCGGAAAC GCTACGAGGA CCTGCTAACC AGGCTGCGGG CCAACCAGAG
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      CTGGGAAGAT TCGAACACCG ACCTCGTCCC GGCCCCTGCA GTCCGGATAC TCACGCCAGA
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      AGTGCGGCTG GGATCCGGCG GCCACCTGCA CCTGCGTATC TCTCGGGCCG CCCTTCCCGA
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      GGGGCTCCCC GAGGCCTCCC GCCTTCACCG GGCTCTGTTC CGGCTGTCCC CGACGGCGTC
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      AAGGTCGTGG GACGTGACAC GACCGCTGCG GCGTCAGCTC AGCCTTGCAA GACCCCAAGC
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      GCCCGCGCTG CACCTGCGAC TGTCGCCGCC GCCGTCGCAG TCGGACCAAC TGCTGGCAGA
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      ATCTTCGTCC GCACGGCCCC AGCTGGAGTT GCACTTGCGG CCGCAAGCCG CCAGGGGGCG
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      CCGCAGAGCG CGTGCGCGCA ACGGGGACGA CTGTCCGCTC GGGCCCGGGC GTTGCTGCCG
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      CATGCACGCG CAGATCAAGA CGAGCCTGCA CCGCCTGAAG CCCGACACGG AGCCAGCGCC
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      CTGCTGCGTG CCCGCCAGCT ACAATCCCAT GGTGCTCATT CAAAAGACCG ACACCGGGGT
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      GTCGCTCCAG ACCTATGATG ACTTGTTAGC CAAAGACTGC CACTGCATAT GAGCAGTCCT
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      GGTCCTTCCA CTGTGCACCT GCGCGGGGA GGCGACCTCA GTTGTCCTGC CCTGTGGAAT
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      GGGCTCAAGG TTCCTGAGAC ACCCGATTCC TGCCCAAACA GCTGTATTTA TATAAGTCTG
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      TTATTTATTA TTAATTTATT GGGGTGACCT TCTTGGGGAC TCGGGGGCTG GTCTGATGGA
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      AAAA
      Seq ID NO: 191 Protein sequence:
      Protein Accession #: NP_004855
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      EDLLTRIRAN QSWEDSNTDL VPAPAVRILT PEVRLGSGGH LHLRISRAAL PEGLPEASRL
HRALFRLSPT ASRSWDVTRP LRRQLSLARP QAPALHLRLS PPPSQSDQLL AESSSARPQL
                                                                                120
50
                                                                                180
      ELHLRPQAAR GRRRARARNG DDCPLGPGRC CRLHTVRASL EDLGWADWVL SPREVQVTMC
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      IGACPSQFRA ANMHAQIKTS LHRLKPDTEP APCCVPASYN PMVLIQKTDT GVSLQTYDDL
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      LAKDCHCI
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      Nucleic Acid Accession #: XM 061731.1
      Coding sequence: 1-567 (underlined sequences correspond to start and stop codons)
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                                          31
                                                                  51
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      GTAAAAGAAC CTTTCTCTCT GATTGGACTT GACACACAGA AGGATCTCAG TAAAGATTTG
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      CTGTTGTTGA TGTCCACAGA CACTGGCAAG GACAGGTTTA CCAACATACT GCTGTCACAC
                                                                                240
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      TCCCTCCAA TGTGCACCAA ATCACGTAAA AATGGGGATA ATGACTCCCC TGCCTTCACA
                                                                                300
      TGGGGTGGCA AAGACACCAG GAGCAATACT GATCTTCCTA TCAGAGACCC TGGGGGCAAG
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      AGTCTTTCAC TCACCAAACA TTCCCACAAG CCTGTCCCTG AGCATCAGTG TGACCAGAGA GAGGTCTTCC AGCCACTTTC AGAGCCAGGT GTAGAAGCAG AGATGGAAGT GTTCGCTGAT
                                                                                420
                                                                                480
      GCTGGATGGT GGATTTATCA GAGCTGTCAG GTTCCTTCCT CAACCCTTGC AAGAAAGAAG
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      ATGGTTTATT CTAAAGAAAC TGAGTGA
      Seq ID NO: 193 Protein sequence:
      protein Accession #: XP 061731.1
75
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MRKGNEGENT EEGRLAQLAQ RKFLKEDGIT LHISLCLSIA VKEPFSLIGL DTOKDLSKDL

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LLLMSTDTGK DRFTNILLSH SPPMCTKSRK NGDNDSPAFT WGGKDTRSNT DLPIRDPGGK
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      SLSLTKHSHK PVPEHQCDQR EVFQPLSEPG VEAEMEVFAD AGWWIYQSCQ VPSSTLARKK
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      TTCTCTTCTC CGCCATGGAA TTCTGCTCCG TGCTTTTAGC CCTCCTGAGC CAAAGAAACC
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      CCAGACAACA GATGCCCATA CGCAGCGTAT AGCAGTAACT CCCCAGCTCG GTTTCTGTGC
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      CTCATATTCT GTTTACACAT CTTGAAAGGC GCTCAGTAGT TCTCTTACTA AACAACCACT
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      GGGGGCCAAA GTGAGCGAAA CCATCCGGAA GGGCTTGATT GACGTGGAGA TGTACAACTC
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      GACTCAAGGG CTACTGATGG CCGGCTCAGT CAGTGCTATG TTTGGTTCTG CTGTGTGGCA
      ACTCGTGGCT TCGTTTTTGA AGCTCCCTAT TTCTGGAACC CATTGTATTG TTGGTGCAAC
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      AATTGTGATG TCTTGGTTCG TGTCCCCACT GCTTTCTGGA ATTATGTCTG GAATTTTATT
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      CTTCCTGGTT CGTGCATTCA TCCTCCATAA GGCAGATCCA GTTCCTAATG GTTTGCGAGC
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      TTTGCCAGTT TTCTATGCCT GCACAGTTGG AATAAACCTC TTTTCCATCA TGTATACTGG
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      CTTGAAAGAA GACCATGAAG AAACAAAGTT GTCTGTTGGT GATATTGAAA ACAAGCATCC
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      TGTTTCTGAG GTAGGGCCTG CCACTGTGCC CCTCCAGGCT GTGGTGGAGG AGAGAACAGT
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      CTCATTCAAA CTTGGAGATT TGGAGGAAGC TCCAGAGAGA GAGAGGCTTC CCAGCGTGGA
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      GTATCACACC GTGCATAAGG ATTCCGGCCT GTACAAAGAG CTACTCCATA AATTACATCT
      TGCCAAGGTG GGAGATTGCA TGGGAGACTC CGGTGACAAA CCCTTAAGGC GCAATAATAG
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      ATCTGAGATA GACATGAGTG TCAAGGCAGC GATGGGTCTA GGTGACAGAA AAGGAAGTAA
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      CAGAATG<u>TGA</u> AGCTGTTTGA GATTAAAATT TGTGTCAATG TTTGGGACCA TCTTAGGTAT
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      TCCTGCTCCC CTGAAGAATG ATTACAGTGT TAACAGAAGA CTGACAAGAG TCTTTTTATT
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     ACATGCACAG GGATTTAACA ACAAAAATAT AACTACAACT TCCCTTGTAG TCTCTTATAT
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     QACILASIFE TVGSVLLGAK VSETIRKGLI DVEMYNSTQG LLMAGSVSAM FGSAVWQLVA
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     SPLKLPISGT HCIVGATIGP SLVAKGQBGV KWSELIKIVM SWFVSPLLSG IMSGILFFLV
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     RAFILHKADP VPNGLRALPV FYACTVGINL FSIMYTGAPL LGFDKLPLWG TILISVGCAV
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      FCALIVWFFV CPRMKRKIER BIKCSPSESP LMEKKNSLKE DHEETKLSVG DIENKHPVSE
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360

VGPATVPLQA VVEERTVSFK LGDLEEAPER ERLPSVDLKE ETSIDSTVNG AVQLPNGNLV

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      DMSVKAAMGL GDRKGSNGSL BEWYDQDKPE VSLLFQFLQI LTACFGSFAH GGNDVSNAIG
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      PLVALYLVYD TGDVSSKVAT PIWLLLYGGV GICVGLWVWG RRVIQTMGKD LTPITPSSGF
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      GAGCGAGCCC CTCCCCGGCT CCAGCCCGGT CCGGGGCCGC GCCGGACCCC AGCCCGCCGT
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      CCAGCGCTGG CGGTGCAACT GCGGCCGCGC GGTGGAGGGG AGGTGGCCCC GGTCCGCCGA
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      CAGGAGAAGC AGCGTGGCCT GCACAGCGAG CTGGGAGAGT CCAGTCTCAT CCTGAAAGCA
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      AACACAGTAT TGCTCAGACA CGACAACATC CTAGGCTTCA TCGCCTCAGA CATGACCTCC
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      CTATCTGGGT AGAGGTAGTG TGAGTGTGGT GTGTGCTGGG GATGGGCAGC TGCGCCTGCC
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      LILGPVLALL ALVALGVLGL WHVRRRQEKQ RGLHSELGES SLILKASEQG DTMLGDLLDS
      DCTTGSGSGL PFLVQRTVAR QVALVECVGK GRYGEVWRGL WHGESVAVKI FSSRDEQSWF
                                                                            240
      RETEIYNTVL LRHDNILGFI ASDMTSRNSS TOLWLITHYH EHGSLYDFLO ROTLEPHLAL
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      RLAVSAACGL AHLHVEIFGT QGKPAIAHRD FKSRNVLVKS NLQCCIADLG LAVMHSQGSD
YLDIGNNPRV GTKRYMAPEV LDEQIRTDCF BSYKWTDIWA FGLVLWEIAR RTIVNGIVED
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      TTTGTGTGAT TTTGCTAAAA TGCATCACCA ACAGCGAATG GCTGCCTTAG GGACGGACAA
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      AGAGCTGAGT GATTTACTGG ATTTCAGTGC GATGTTTTCA CCTCCTGTGA GCAGTGGGAA
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      AAATGGACCA ACTTCTTTGG CAAGTGGACA TTTTACTGGC TCAAATGTAG AAGACAGAAG
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      CHQQSLLGGD MDMGNPGTLS PTKPGSQYYQ YSSNNPRRRP LHSSAMEVQT KKVRKVPPGL
                                                                           180
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      PSSVYAPSAS TADYNRDSPG YPSSKPATST FPSSFFMQDG HHSSDPWSSS SGMNQPGYAG
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      MLGNSSHIPQ SSSYCSLHPH ERLSYPSHSS ADINSSLPPM STFHRSGTNH YSTSSCTPPA
      NGTDSIMANR GSGAAGSSQT GDALGKALAS IYSPDHTNNS FSSNPSTPVG SPPSLSAGTA
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      SATSPDLNPP QDPYRGMPPG LQGQSVSSGS SEIKSDDEGD ENLODTKSSE DKKLDDDKKD
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      IKSITSNNDD EDLTPEOKAE REKERRMANN ARERLRVRDI NEAFKELGRM VQLHLKSDKP
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      QTKLLILHQA VAVILSLEQQ VRERNLNPKA ACLKRREEEK VSSEPPPLSL AGPHPGMGDA
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      AACCTACTGG CCTATCTAAA GCACCTCAAA GGGCAAAACG AGGCAGCCCT GGAATGCTTA
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      GAAAGAGCGA AGGTGTGCTT TGAGAAGGCT CTGGAAAAGA AGCCAAAGAA CCCAGAATTC
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      ACCICIGGAC TGGCAATAGC AAGCTACCGI CTGGACAACT GGCCACCATC TCAGAACGCC
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      CTGGCTCTGA AGCTTCATAA GATGCGTGAA GAAGGTGAAG AGGAAGGTGA AGGAGAGAAG
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      AMETITTATC GAAGAAAGA TGAGCCAGAC AAAGCGATTG AACTGCTTAA AAAGGCTTTA
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      GAATACATAC CAAACAATGC CTACCTGCAT TGCCAAATTG GGTGCTGCTA TAGGGCAAAA
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      CGTGTCTGTT CCATTCTTGC CAGCCTCCAT GCTCTAGCAG ATCAGTATGA AGAAGCAGAG
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      GTCTTGGCAT TCCTTCAGGA GCTGAATGAA AAAATGCAAC AAGCAGATGA AGACTCTGAG
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      ATAGAGATGT GGTGCCCACT AGGCTACTGC TGAAAGGGAG CTGAAATTCC TCCACAAGTT
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      CACTGGACAG GGTTATGTTA AACCTGAATT GCTGGGTCTT AAAAGAGCCC AAGGAGTTCT
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      CTCTTTGCGG AACTTC
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                                                                           120
      KHVCEKFSSP YRIESPELDC EEGWTRLKCG GNQNERAKVC FEKALEKKPK NPEFTSGLAI
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      ASYRLDNWPP SQNAIDPLRQ AIRLNPDNQY LKVLLALKLH KMREEGEEG EGEKLVEEAL
                                                                           240
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      EKAPGVTDVL RSAAKFYRRK DEPDKAIELL KKALEYIPNN AYLHCQIGCC YRAKVFQVMN
                                                                           300
      LRENGMYGKR KLLELIGHAV AHLKKADEAN DNLFRVCSIL ASLHALADQY EDAEYYFQKE
                                                                           360
      FSKELTPVAK QLLHLRYGNF QLYQMKCEDK AIHHFIEGVK INQKSREKEK MKDKLQKIAK
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      MRLSKNGADS EALHVLAFLQ ELNEKMQQAD EDSERGLESG SLIPSASSWN GE
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                                                                           300
      GACTTGATCA GGCTCTGCCC TGTCTGACAG AACTCATTCT CACCAATAAT AGTCTCGTGG
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      AACTGGGTGA TCTGGACCCT CTGGCATCTC TCAAATCGCT GACTTACCTA AGTATCCTAA
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      ATCCAGGTGC TGGTTTGCCA ACTGACAAAA AGAGAGGTGG GCCATCTCCA GGGGATGTAG
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      ATAATAATAG GCCCTCTTGG AACAAGTCTT GCTTTTCGAA CATGGTATAA TAGCCTTGTT
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      TGTGTTAGCA AAGTGGAATC TATCAGCATT GTTGAAATGC TTAAGACTGC TGCTGATAAT
                                                                           960
      TTTGTAATAT AAGTTTTGAA ATCTAAATGT CAATTTTCTA CAAATTATAA AAATAAACTC
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      Seq ID NO: 203 Protein sequence:
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      LRNPVTNKKH YRLYVIYKVP QVRVLDFQKV KLKERQBAEK MFKGKRGAQL AKDIARRSKT
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      FNPGAGLPTD KKRGGPSPGD VEAIKNAIAN ASTLAEVERL KGLLQSGQIP GRERRSGPTD
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      TGGGAGGAGC CTGAGGTGGA CCTCGCGTCC TGAGTCTGGA AGGCACCTAT GGGGACCTGC
                                                                           180
      TGGGGTGATA TCTCAGAAAA TGTGAGAGTA GAAGTTCCCA ATACAGACTG CAGCCTACCT
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      ATCCATCCAG TTGGTTGGTG TGCAGCCAGC GGAAAACCTC TTGTTCCTCC TAGAACTATT
      CAGCATAAAT ATACAAACTG GAAAGCTTTT CTAGTGAAAC GACTTACTGG TGCCAAAACA
                                                                             480
      CTGCCTCCTG ATTTCTCCCA AAAGGTTTCA GAGAGTATGC AGTATCCTTT CAAACCTTGC
                                                                             540
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                                                                           1320
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20
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                                                                           1680
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      CTGCCGGGCG TGGTGGCACA CGCCTGAAGT CCCAGCTACT CAGGAGGCTG AGGCAAGAGA
                                                                           2460
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      CTGGGCAACA CAGCAAGACT CTGTCTCAAA AAAAAAAAA AAAA
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      FKPCMRVEVV DKRHLCRTRV AVVESVIGGR LRLVYEESED RTDDFWCHMH SPLIHHIGWS
RSIGHRFKRS DITKKQDGHF DTPPHLFAKV KEVDQSGEWF KEGMKLEAID PLNLSTICVA
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                                                                            240
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      TIRKVLADGF LMIGIDGSEA ADGSDWFCYH ATSPSIFPVG FCEINMIELT PPRGYTKLPF
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TATAAGAGAA ATCAAATATA TACCTACATC GGCTCCATCC TGGCCTCCGT GAACCCCTAC
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      MAYIMLCRKT SQOEHVYEAA RAHAREANDS GETMRVAIFA SGCSSDEPTS QNLGNNYSDE
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Seq ID NO: 229 Protein sequence:

## Protein Accession #: NP_006024.1

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It is understood that the examples described above in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All publications, sequences of accession numbers, and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference.

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## WHAT IS CLAIMED IS:

1 1. A method of detecting an angiogenesis-associated transcript in a cell in 2 a patient, the method comprising contacting a biological sample from the patient with a 3 polynucleotide that selectively hybridized to a sequence at least 80% identical to a sequence 4 as shown in Tables 1-8. 1 2. The method of claim 1, wherein the biological sample is a tissue 2 sample. 1 3. The method of claim 1, wherein the biological sample comprises 2 isolated nucleic acids. 1 4. The method of claim 3, wherein the nucleic acids are mRNA. 1 5. The method of claim 3, further comprising the step of amplifying 2 nucleic acids before the step of contacting the biological sample with the polynucleotide. 1 6. The method of claim 1, wherein the polynucleotide comprises a 2 sequence as shown in Tables 1-8. 1 7. The method of claim 1, wherein the polynucleotide is labeled. 1 8. The method of claim 7, wherein the label is a fluorescent label. 1 9. The method of claim 1, wherein the polynucleotide is immobilized on 2 a solid surface. 1 10. The method of claim 1, wherein the patient is undergoing a therapeutic 2 regimen to treat a disease associated with angiongenesis. 1 11. The method of claim 1, wherein the patient is suspected of having 2 cancer. 1 12. An isolated nucleic acid molecule consisting of a polynucleotide 2 sequence as shown in Tables 1-8. 1 13. The nucleic acid molecule of claim 12, which is labeled.

1

14.

The nucleic acid of claim 13, wherein the label is a fluorescent label

1	13.	An expression vector comprising the nucleic acid of claim 12.
1	16.	A host cell comprising the expression vector of claim 15.
1 2	17. having polynucleoti	An isolated polypeptide which is encoded by a nucleic acid molecule de sequence as shown in Tables 1-8
1	18.	An antibody that specifically binds a polypeptide of claim 17.
.1	19. component.	The antibody of claim 18, further conjugated or fused to an effector
1 2	20. fluorescent label.	The antibody of claim 19, wherein the effector component is a
1 2	21.	The antibody of claim 19, wherein the effector component is a
1	22.	The antibody of claim 19, which is an antibody fragment.
1	23.	The antibody of claim 19, which is a humanized antibody
1	24. sample from a patien	A method of detecting a cell undergoing angiogenesis in a biological at, the method comprising contacting the biological sample with an
3	antibody of claim 18	
1	25. fused to an effector	The method of claim 24, wherein the antibody is further conjugated or component.
1 2	26. fluorescent label.	The method of claim 25, wherein the effector component is a
1	27.	The method of detecting antibodies specific to angiogenesis in a
2	-	comprising contacting a biological sample from the patient with a
3	polypeptide which is	s encoded by a nucleotide sequence of Tables 1-8.